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(54) Title: CYCLIN DEPENDENT KINASE (CDK)4 INHIBITORS AND THEIR USE FOR TREATING CANCER

(57) Abstract

Certain derivatives of acridones and benzothiadiazines have been found to have anti-cancer properties by virtue of their specific inhibition of the cyclin D dependant kinase CDK4. These molecules inhibit CDK4 activity more than they inhibit the activity of other such kinases (e.g. CDC2 and CDK2). This specificity results in an improved therapeutic index when used as drugs to treat susceptible cancers.

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CYCLIN DEPENDENT KINASE (CDK)4 INHIBITORS AND THEIR USE FOR TREATING CANCER

I. FIELD OF THE INVENTION

The present invention concerns compounds that inhibit cyclin-dependent kinases, particularly the cyclin-dependent kinase CDK4, and methods for treating cancers using such compounds.

II. BACKGROUND OF THE INVENTION

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In a normal cell CDK4:cyclin D kinase holoenzyme phosphorylates the retinoblastoma protein (Rb) to form hyperphosphorylated retinoblastomaphosphate (Rb-p). The hyperphosphorylation of retinoblastoma protein results in the release of Rb-p associated transcription factors that allow cell cycle progression beyond the G1 checkpoint, thereby promoting cell proliferation (Schrr et al., U.S. Patent No. 5,723,313, (1998)).

The p16 gene (also known as CDKN2, MST1, and CDK4I) encodes the protein p16^{INK4A}, which inhibits the cyclin-dependent kinase (CDK)4:cyclin D complex (Serrano, et al., Nature 366: 704-7 (1993)). Defects in the p16/CDK4:cyclinD/Rb pathway may lead to tumor formation. Genetic alteration or over expression of CDK4 and CyclinD1 has been observed in various tumor cell types. In addition, alterations of p16 have been described in various histologic types of human cancers including retinoblastoma, astrocytoma, melanoma, leukemia, breast cancer, head and neck squamous cell carcinoma, malignant mesothelioma, and lung cancer (Kamb et al., Science 264: 436-40 (1994); Noborie et al., Nature 368: 753-56 (1994); Walker et al., Cancer Res. 55: 20-3 (1995) and Nakagawa et al., Oncogene 11: 1843-51 (1995)).

Acridones and Benzothiadiazines

Acridones and benzothiadiazines (BTDs) are classes of known cyclic aryl compounds. Certain known acridones or BTDs have pharmacological effects. For example, BTDs have been investigated as diuretics (See de Tullio et al., *J. Med. Chem*). Fajans and Floyd (*Ann. Rev. Med.* 30:313-329, 1982)

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disclose the use of "diuretic benzothiadiazine, e.g. trichlormethiazide" as a hyperglycemic in the treatment of insulinomas. Fajans and Floyd, however, do not teach the use of BTDs to affect cancers directly. The prior art, as understood, does not appear to teach the use of BTDs for their direct antineoplastic effect in the specific inhibition of CDK4 dependent tumors.

Particular acridones and acridines are known. For example, (C₁₈H₁₉N₃O₂-HCl) has been mentioned in a paper concerned with the anti-tumor activity of linear tri-cyclic carboxamides (Palmer et al., J. Med. Chem (US) 31 (4) pgs.707-721, 1988). Interestingly, the Palmer et al. paper states that this compound is "inactive" (page 711, column 1, paragraph 3).

The basic *thio*acridone ring structure was described in DeLeenheer et al., *J. Pharm. Sci.* 60:1238-1239, 1971, and is shown below.

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1-nitro-9-acridone, 1-nitro-10-(3-N,N-dimethylaminopropryl)-9-acridone, 1-amino-2,4-diethylthio-9-acridone and a number of acridine derivatives have been disclosed by Weltrowski et al. (*Pol. J. Chem Technol.* 56:77-82, 1982). This paper, however, deals exclusively with the synthesis of nitroacridines and does not discuss any biological activity or mechanism of biological action. But, the title of the Weltrowski article refers to tumor inhibition, and the footnote states that the work was supported by the Polish National Cancer Program.

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III. SUMMARY OF THE INVENTION

The present invention concerns acridones, benzothiadiazines and derivatives thereof that are useful for treating cancers. The invention also concerns methods for using these compounds as CDK4 inhibitors to treat cancers.

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There are a number of dreadful and relatively common cancers that have been shown to involve alterations in p16. These cancers include lung cancer, breast cancer, melanoma, leukemia, retinoblastoma, astrocytoma, head and neck squamous cell carcinoma and malignant mesothelioma. Expression of normal p16 protein in tumor cells with alterations of p16 results in restoration of cell-cycle regulation, decreased cell growth and decreased tumorigenicity *in vivo*. Because the only known function of p16 is inhibition of CDK4 kinase activity, cancers with alterations of p16, including those listed above, are likely to be sensitive to CDK4 inhibitors. Prior inhibitors of cyclin-dependent kinases, such as flavopiridole, staurosporin, and UCN-01, inhibit CDC2 and CDK2 as well as the intended target, CDK4. This lack of specificity produces pathological side effects, such as bone marrow and gastrointestinal toxicities, and limits their clinical application.

As a result, there is a need for drugs for treating CDK4 sensitive neoplasms that minimize toxic side effects caused by concomitant inhibition of CDC2 and CDK2. The compounds claimed in this application inhibit CDK4 to a far greater extent than CDC2 or CDK2 and therefore satisfy this need.

One example of a novel compound of the present invention is 3-amino-9- thio(10H)-acridone. This compound and others can be used to form therapeutic compositions. One embodiment of such a composition comprises a therapeutically effective amount of a compound selected from the group consisting of a benzothiadiazine, a thioacridone, or mixtures thereof. The compound has an IC₅₀ for CDK4 of less than about 10 μ M, preferably from about 1 μ M to about 7 μ M, an IC₅₀ for CDC2 of greater than about 60 μ M, preferably greater than about 100 μ M, an IC₅₀ for CDK2/A of greater than about 100 μ M, an IC₅₀ for CDK2/E of greater than about 80 μ M, and preferably greater than about 100 μ M.

The specificity of the compounds for inhibiting CDK4 can be expressed as a ratio of the IC₅₀ values for other enzymes relative to CDK4. Such compositions typically comprise a compound selected from the group consisting of a benzothiadiazine, a thioacridone, or mixtures thereof, the compound having an IC₅₀ ratio for CDC2:CDK4 of greater than about 8.5, typically greater than about 20, preferably greater than about 60; an IC₅₀ ratio for CDK2/A:CDK4 of

greater than about 14, typically greater than about 20, and preferably greater than about 60; and an IC_{50} ratio for CDC2/E:CDK4 of greater than about 11.5, typically greater than about 20, and preferably greater than about 60.

The invention also provides a composition comprising an effective amount of a compound according to Formula 1

where m is 0 or 1, n=m, R_1-R_4 are independently selected from the group consisting of H, $-NH_2$ and lower alkoxy, where with m=1 one of R_1-R_4 is an amine bonded to R' to form an arylamide,

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where R and R_1 are independently carbon or nitrogen, where if R_1 =carbon X is hydrogen, halogen, aryl or alkoxy, and R_2 is selected from the group consisting of lower alkyl and aryl amino. The composition also can comprise mixtures of compounds satisfying Formula 1 and/or Formula 2. The composition can further include, without limitation, additives selected from the group consisting of carriers, diluents, excipients, diagnostics, direct compression buffers, buffers, stabilizers, fillers, disintegrates, flavors, colors, and mixtures thereof.

A method for inhibiting the growth of living cells also is described.

The method comprises providing a compound selected from the group consisting

of a benzothiadiazine, a thioacridone, or mixtures thereof, as described above. An effective amount of the compound, a mixture of compounds, or a composition comprising the compound or mixture of compounds, is administered to a subject to inhibit the growth of living cells.

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IV. BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1(A)-1(I) are dose-response curves showing the effect of Compound 5 on various cancer cell lines in culture.

FIG. 2 shows mean plots of data from FIGS. 1A-1I, wherein the left-hand mean plot is of GI_{50} data, the middle mean plot is of TGI data, and the right-hand mean plot is of LC_{50} data.

FIGS. 3(A)-3(I) are dose-response curves showing the effect of Compound 7 on various cancer cell lines in culture.

FIG. 4 shows mean plots of data from FIGS. 3A-3I, wherein the left-hand mean plot is of GI_{50} data, the middle mean plot is of TGI data, and the right-hand mean plot is of LC_{50} data.

FIGS. 5(A)-5(I) are dose-response curves showing the effect of Compound 8 on various cancer cell lines in culture.

FIG. 6 shows mean plots of data from FIGS. 5A-5I, wherein the left-hand mean plot is of GI_{50} data, the middle mean plot is of TGI data, and the right-hand mean plot is of LC_{50} data.

FIGS. 7(A)-7(I) are dose-response curves showing the effect of Compound 4 on various cancer cell lines in culture.

FIG. 8 shows mean plots of data from FIGS. 7A-7I, wherein the left-hand mean plot is of GI_{50} data, the middle mean plot is of TGI data, and the right-hand mean plot is of LC_{50} data.

FIGS. 9(A)-9(I) are dose-response curves showing the effect of Compound 6 on various cancer cell lines in culture.

FIG. 10 shows mean plots of data from FIGS. 9A-9I, wherein the left-hand mean plot is of GI_{50} data, the middle mean plot is of TGI data, and the right-hand mean plot is of LC_{50} data.

FIGS. 11(A)-11(I) are dose-response curves showing the effect of

Acres 1

Compound 3 on various cancer cell lines in culture.

FIG. 12 shows mean plots of data from FIGS. 11A-11I, wherein the left-hand mean plot is of GI_{50} data, the middle mean plot is of TGI data, and the right-hand mean plot is of LC_{50} data.

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V. DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS DEFINITIONS

Particular terms and phrases used herein typically have the meanings set forth below. These definitions are provided solely for convenience and should not be interpreted to limit the invention to a scope less than that known to a person of ordinary skill in the art.

"3-ATA" means 3-amino-9-thio(10H)-acridone.

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"BTD" means benzothiadiazine.

"Neoplasm" and "cancer" both refer to any cell or tissue wherein growth and cell division have become uncoupled from the normal regulatory constraints of the cell cycle to produce a pathological state.

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"Tumor" is any neoplasm and includes both solid and non-solid neoplasms.

"Inhibitory concentration" or " IC_{50} " means the drug concentration at 50% inhibition of kinase activity (μM).

"Therapeutically effective anti-neoplastic amount" means an amount sufficient to prevent advancement, or to cause regression of, a neoplasm.

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"CDK4" and "CDK4/A" refer to the CDK4:cyclin D1 kinase holoenzyme.

"CDK4 inhibitor" refers to compounds that inhibit the kinase activity of CDK4.

"CDK4 inhibition" refers to inhibition of the kinase activity of CDK4.

5 "CDK2", when used alone, refers to both CDK2:Cyclin A and to CDK2:Cyclin E

"CDC2" and "CDC2/A" refer to CDC2:Cyclin A holoenzyme.

"CDK2/A" refers to CDK:Cyclin A holoenzyme.

"CDK2/E" refers to CDK2:Cyclin E holoenzyme.

"Cancers specifically inhibited by CDK4 inhibitors" means all neoplastically transformed cells and tissues, the growth and/or cell cycle of which is affected by a CDK4 inhibitor.

A cell "susceptible to CDK4 inhibitors" or "susceptible to CDK4 inhibition" is a cell for which CDK4 inhibitors alter growth or cell cycle.

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"Specific inhibition" or "specific inhibitory activity" of the compounds of the invention means that the compounds inhibit CDK4 to a greater extent than they inhibit CDC2 or CDK2.

"Lower alkyl" means a single-bonded branched or unbranched hydrocarbon chain having from about one to about ten carbon atoms, including all position and stereoisomers.

COMPOUNDS

Compounds of the present invention satisfy either Formula 1 (acridone-like structures) or Formula 2 (benzothiadiazine-like structures) below.

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FORMULA 1

FORMULA 2

N H R₁ - X

With reference to Formula 1, m is 0 or 1, and n = m. R_1 - R_4 are independently selected from the group consisting of H, -NH₂ and lower alkoxy. With m=1, at least one of R_1 - R_4 is an amine and R' is bonded to the amine to form an arylamide.

With reference to Formula 2, R and R_1 are independently carbon or nitrogen. If R_1 =carbon X is hydrogen or halogen. R_2 is selected from the group consisting of lower alkyl and aryl amino.

Compounds according to both Formula 1 and 2 show specific inhibitory activity against CDK4. This inhibition may be due to inhibition of formation of the CDK4:cyclinD kinase holoenzyme or to competitive binding of the inhibitor with the kinase substrate or to ATP-dependent competitive effects or some other interaction.

Structural formulas for particular compounds of the invention are provided below as Compounds 1-6.

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COMPOUND 1

S NH₂

3-Amino-10H-acridine-9-thione

COMPOUND 2

S OCH,

1,4-Dimethoxy-10H-acridine-9-thione

COMPOUND 3

S H

2,2'-Biphenyldiamine, bis[N,N'-[3-(amidonmethylamino)-10H-acridine-9-thione]]

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COMPOUND 4

4-(4-Fluorobenzylamino)-1,2,3-benzothiadiazine-1,1-dioxide

10 COMPOUND 5

3-Chloro-4-methyl-4H-benzo[e][1,2,4]thiadiazine 1,1-dioxide

COMPOUND 6

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3-Chloro-4-ethyl-4H-benzo[e][1,2,4]thiadiazine 1,1-dioxide

SYNTHESIS OF COMPOUNDS

The compounds of the invention were obtained from and are
maintained at the Drug Synthesis and Chemistry Branch, National Cancer
Institute. Syntheses of related compounds are known in the literature. For
example, the following references described the syntheses of certain related

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compounds: Pascal de Tullio et al., "3- and 4- Substituted 4*H*-Pyrido[4,3-e]-1,2,4-thiadiazine 1,1-Dioxides as Potassium Channel Openers: Synthesis, Pharmacological Evaluation, and Structure--Activity Relationships," *J. Med. Chem.*, Vol. 39, pp. 937-948 (1996); Bernard A. Dumaitre et al., U.S. Patent No. 5,604,237; Hamprecht et al., U.S. Patent No. 4,075,004; Magatti U.S. Patent No. 4,468,396; Brian D. Palmer et al., "Potential Antitumor Agents. 54. Chromophore Requirements for in Vivo Antitumor Activity Among the General Class of Linear Tricyclic Carboxamides," *J. Med. Chem.*, Vol. 31, pp. 707-712 (1988); N. Dodic et al., "Synthesis and Activity Against Multidrug Resistance in Chinese Hamster Ovary Cells of New Acridone-4-Carboxamides," *J. Med. Chem.*, Vol. 38, pp. 2418-2426 (1995); Marek Welt4rowski et al., "Research on Tumour Inhibiting Compounds, Part LXX, Reactions of 1-Nitroacridines with Ethanethiol," *Polish Journal of Chemistry*, pp. 77-82 (1982).

15 **COMPOSITIONS**

Compounds satisfying either Formula 1 or 2 above may be formulated as pharmacological compositions containing a therapeutically effective antineoplastic amount of the compound(s). Such compositions may further comprise, without limitation, inert carriers, diluents, excipients, diagnostics, direct compression buffers, buffers, stabilizers, fillers, disintegrates, flavors, colors, other materials conventionally used in the formulation of pharmacological compositions and mixtures thereof.

METHOD

The method of the present invention comprises administering to a subject a therapeutically effective anti-neoplastic amount of a compound, mixture of compounds, or composition or compositions comprising the compound or compounds, to effect a change in the physiology of a neoplasm. One of ordinary skill in the art will realize that the therapeutically effective anti-neoplastic amount may vary. Anti-tumor agents generally are dosed as mass-per-unit-body surface area of the subject. It currently is believed that a therapeutically effective anti-neoplastic amount of the disclosed compounds may be from about 1 µg to about

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10 g per m² of body surface area, more preferably from about 1 mg to about 900 mg per m² of body surface area. Moreover, it typically is desirable to provide as large a dose as a subject will tolerate.

The compound(s) or compositions may be administered by any number of methods including, but not limited to, intravenously, topically, orally, intramuscularly, subcutaneously, intraperitoneally. Currently, intravenous and oral administration are considered the preferable routes of administration.

BIOLOGICAL METHODS AND RESULTS

Tables 1 and 2 provide IC_{50} data for compounds representative of the present invention. These tables demonstrate that the IC_{50} value of compounds according to the present invention for CDK4 generally is less than about 10 μ M, and preferably is less than about 7 μ M. The best compound, solely in terms of its IC_{50} value for CDK4, is compound 5 with an IC_{50} of 1.1 μ M. But, compounds 7 and 8 also have IC_{50} values of less than 2 μ M, namely 1.4 μ M and 1.7 μ M respectively.

The compounds of the present invention also are quite specific for inhibition of CDK4. This is reflected in the IC_{50} ratios reported in Tables 1 and 2, with the IC_{50} for CDK4 being the denominator in the ratio e.g., $(IC_{50} \text{ CDC2})/(IC_{50} \text{ CDK4})$. Thus, the lower the IC_{50} is for CDK4 and the higher it is for the other complexes, the more specific the compound is for CDK4.

The CDC2/A:CDC4 ratios in Tables 1 and 2 range from about 8 to greater than 72. The best compound with respect to specificity between CDK4 and CDC2 is compound 7, with an IC₅₀ for CDK4 of 1.4 μ M, an IC₅₀ for CDC2 of >100 μ M, and an (IC₅₀ CDC2):(IC₅₀ CDK4) of >71.5.

Compound 3 (3-ATA) has an IC₅₀ for CDK4 of 6.8 μ M, an IC₅₀ for CDC2 of 60 μ M, and an (IC₅₀ CDC2):(IC₅₀ CDK4) of 8.8.

Compound 4 has an IC₅₀ for CDK4 of 2.2 μ M, an IC₅₀ for CDC2 of >100 μ M, and an (IC₅₀ CDC2):(IC₅₀ CDK4) of >45.

Compound 5 has an IC₅₀ for CDK4 of 1.1 μ M, an IC₅₀ for CDC2 of >70 μ M, and an (IC₅₀ CDC2):(IC₅₀ CDK4) of >63.6.

Compound 6 has an IC $_{50}$ for CDK4 of 5.0 μM_{\star} an IC $_{50}$ for CDC2 of

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>100 μ M, and an (IC₅₀ CDC2):(IC₅₀ CDK4) of >71.5.

Compound 8 has an IC₅₀ for CDK4 of 1.7 μ M, an IC₅₀ for CDC2 of >100 μ M, and an (IC₅₀ CDC2):(IC₅₀ CDK4) of >58.8.

 IC_{50} and IC_{50} ratio data for other kinases are summarized in Tables 1 and 2 below.

Compounds satisfying Formulas 1 and 2 have been subjected to biological assays to determine inhibition of the cyclin dependent kinases CDK4, CDC2, CDK2/A and CDK2/E. The experimental procedures for these biological methods and assays are provided below in the Examples. Results of these assays for representative compounds are provided below in Tables 1 and 2.

TABLE 1

	Formula Name	IC ₅₀ value (μM)								
		CDK4/D1	CDC2/A	Ratio CDC2A: CDK4	CDK2/A	Ratio CDK2/A: CDK4	CDK2/E	Ratio CDK2/E: CDK4		
5	Compounds structurally related to 3-ATA									
	Formula 3	6.8	60	8.8	>100	>14.7	80	11.8		
10	Formula 4	2.2	>100	>45	>100	>45	>100	>45		
15	Formula 5	1.1	70	63.6	>100	>91	>100	>91		

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TABLE 2

	Formula Name	IC ₅₀ value (μM)								
	Name	CDK4/D1	CDC2/A	Ratio CDC2A: CDK4	CDK2/A	Ratio CDK2/A: CDK4	CDK2/E	Ratio CDK2/E: CDK4		
5 Compounds structurally related to BTD (NSC645787)										
	Formula 6	5.0	>100	>20	>100	>20	>100	>20		
10	Formula 6	1.4	>100	>71.5	>100	>71.4	>100	>71.4		
15	Formula 7	1.7	>100	>58.8	>100	>58.8	>100	>58.8		

An IC $_{50}$ of 10 μM is generally considered effective for these compounds, but effectiveness should be considered in the light of specificity for CDK4.

EXAMPLES

The following examples are provided to illustrate certain features of the invention and are not meant to limit the invention to any particular embodiment.

Example 1

This example describes in detail how the compounds of the invention were identified and tested to determine their specific inhibitory activity against cyclin dependent kinases. Essentially, the methods of this example include three stages: (1) determining which cell lines contain p16 alterations, (2) determining which drugs are most active against p16 altered cells, and (3) determining the CDK4 kinase inhibitory activity of selected, screened compounds.

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METHODS

Cell lines, compounds, and in vitro sensitivity testing.

Exponentially growing cultures of the nine non-small cell lung, eight melanoma, eight renal, eight breast, seven colon, six brain, six leukemia, six ovarian, and two prostate cancer cell lines from the NCI drug screen panel were used. Compounds were obtained from the Drug Synthesis and Chemistry Branch, National Cancer Institute. *In vitro* antitum or activity of compounds was determined using a sulforhodamine-B assay in the 60 human cancer cell lines of the NCI drug screen panel.

Polymerase chain reaction-single strand conformation polymorphism (PCR-SSCP) and DNA sequence analysis of p16. Approximately 1.5 X 105 tumor cells were washed with PBS, lysed in 100 μl proteinase K solution [200 mg/ml, 50 mM Tris-HC1 (pH8.5), 1 mM EDTA(pH8.0), and 0.5% Tween, 20], and incubated at 50°C for 4 h. One microliter of this lysate was used as template in a 10 µl PCR for each of seven oligonucleotide primer pairs which span the coding region and splice junctions of exons 1 and 2 of p16 twice. Smal-digested (for primer pair 2D) or undigested PCR products were subjected to SSCP. The presence of bands with an abnormal migration pattern was confirmed by repeating PCR-SSCP at least once prior to extraction of the band, cloning into pT7Blue(R) T-vector (Novagen, Madison, WI), and DNA sequence analysis by the dideoxy chain termination method using SequenaseTM (US Biochemical, Cleveland, OH). The presence of intact genomic DNA was confirmed by amplification of a 536-bp fragment of the β-globin gene. The p16 sequence published by Okamoto et al. (GenBank accession number L27211) was used as reference for DNA and amino acid numbering.

Reverse Transcription (RT)-PCR and Southern blot hybridization analyses of p16. Total RNA was isolated from 1 X 10⁶ cells of each cell line using an RNA isolation kit (5' prime 3' prime,Inc., Boulder, CO), RT-PCR was performed for the p16 gene as previously described. PCR products were separated by agarose gel electrophoresis, transferred to a nylon membrane, and hybridized with a 388-bp p16 exon 1 genomic fragment defined by oligonucleotides 2F and 1108R. Expression of the glyceraldehyde-3-phosphate

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(GAPDH) gene was examined to assure the presence of intact mRNA in each sample by addition of a gene-specific oligonucleotide, G3PD-2R (5'-GATACATGACAAGGTGCGGC-3') to the reverse transcriptase reaction followed by 40 cycles of PCR (30 sec at 94°C, 30 sec at 55°C, and 1 min at 72°C using oligonucleotides, G3PD-1F (5'TCGTGGAAGGACTCATGACC-3') and G3PD-1R (5'ACATGGCAACTGTGAGGAGG-3').

Immunoblot analysis. Cells (1 X 10⁷) were washed with PBS, resuspended in 0.4ml of lysis buffer [50 mM Tris-HC1 (pH7.4), 250 mM NaCl, 5 mM EDTA, 0.1% Nonidet P40, 50 mM NaF, and 1 mM PMSF], and centrifuged at 14,000 rpm for 20 min at 4°C. The protein concentration of the supernatant was determined using the Bio-Rad protein assay reagent (Bio-Rad, Hercules, CA). Fifty micrograms of total protein were mixed with an equal volume of 2X sample buffer [125 mM Tris-HC1 (pH 6.8), 20% glycerol, 4% (w/v) SDS, 0.005% bromophenol blue, and 5% 2-mercaptoethanol], loaded on a 14% Tris-glycine gel, and subjected to electrophoresis at 125 V for 90 min in 1X running buffer (25 mM Tris-base, 192 mM glycine, and 0.1% SDS). The separated proteins were transferred to a nitrocellulose membrane at 25 V for 2 h in transfer buffer (12 mM Tris-base, and 96 mM glycine, 20% methanol). After 30 min incubation at room temperature in blocking solution (1X PBS, 5% powdered dry milk, and 1% BSA), the membrane was incubated at 4°C with 1:1000 dilution of polyclonal anti-human p16 antiserum (PharMingen, San Diego, CA) overnight, rinsed 5 times with PBS, incubated with a mixture of 40 µl ¹²⁵ I-Protein A (>30 mCi/mg) in 20 ml blocking solution at 4°C for one hour, washed again with PBS, air dried for 15 min, and subjected to autoradiography.

COMPARE analysis. The COMPARE algorithm was performed. For the identification of agents with differential activity, "G150" values of 0 and 1 were used for p16-normal and for p16-altered cell lines, respectively. p16-altered cell lines were those with biallelic deletion, intragenic mutation, or transcriptional suppression of p16 and p16-normal cell lines were those without these abnormalities. Pearson correlation coefficients were calculated by the SAS procedure PROC CORR (SAS Institute Inc., Cary, NC).

GST fusion proteins. Full length p16 cDNA from cell lines

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10 In vitro kinase assay. Seventy-two hours after infection of 1 X 10⁷ Sf9 cells with baculovirus containing a human CDK gene and/or a cyclin gene, cells were lysed in 250 µl of lysis buffer [50 mM HEPES (pH 7.5), 10 mM MgCl₂, 1 mM DTT, 5 ig/ml of aprotinin, 5 µg/ml of leupeptin, 0.1 mM NaF. 0.2 mM phenylmethylsulfonyl fluoride (PMSF), and 0.1 mM sodium orthovanadate], centrifuged, and lysates stored at -70°C. Five microliters of 15 CDK:cyclin lysate were mixed with test compounds in 40 µl of kinase buffer (200 mM Tris-HCl, pH 8.0, 100 mM MgCl₂, 10 mM EGTA) and incubated at 30°C for 30 min. About 400 ng of purified GST-Rb fusion protein and 5 μCi of γ-[³²P]ATP were added to the mixture and incubated at 30°C for 15 min. 20 Reactions were stopped by the addition of 250 µ1 of IP buffer (50 mM Tris-HC1, pH 8.0; 150 mM NaC1, 0.5% NP-40) and 15 μ 1 glutathione sepharose. After one hour incubation at 4°C, sepharose beads were washed four times with IP buffer, mixed with 18 μ1 of 2X sample buffer and electrophoresed on an 8% Tris-glycine gel (Novex, San Diego, CA) at 125 V for 90 min. Equal recovery of GST-Rb fusion protein was confirmed by Coomassie blue staining prior to 25 autoradiography.

CDK4 binding assay. Sf9 cells (1 X 10⁷) were co-infected with baculovirus containing a cloned human CDK4 gene and/or a cyclin D1 gene in 12.5 ml of Grace's insect medium (Paragon, Baltimore, MD) containing 10% FBS. After 40 h, cells were washed and placed in 5 ml of methionine-free medium containing 200 μCi/ml of [35S]methionine (1000 Ci/mmole) for 4 h, followed by lysis in 250 μl. Cleared cell lysate (10 μl) was incubated with 400

ng of wildtype or mutant GST-p16 fusion proteins using the same conditions as the *in vitro* kinase assay. After a 30 min incubation, GST-p16 fusion protein was separated using glutathione sepharose according to manufacturer's recommendations, and electrophoresed on a 14% Tris-glycine gel (Novex, San Diego, CA). The gel was stained using Coomassie blue, dried, and autoradiography was performed. Equal recovery of GST-p16 fusion protein was confirmed by Coomassie blue staining. To test the effect of compounds on p16 binding to CDK4, 100 μ M of each compound was incubated with CDK4:cyclin D1 lysate for 30 min prior to adding GST-p16 fusion protein.

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RESULTS

Characterization of the p16 status of the cell lines of the NCI drug screen panel. To detect genetic alternations of p16 in the 60 cell lines of the NCI drug screen panel, polymerase chain reaction-single strand conformation polymorphism (PCR-SSCP) analysis was performed for exons 1 and 2 of the p16 gene using genomic DNA. Exon 3, which encodes only four amino acids, was not examined as mutations limited to exon 3 have not been described. Among the 60 cell lines, 29 cell lines were found to lack amplifiable genomic sequences of one or both exons, indicative of a biallelic deletion involving p16. The presence of amplifiable genomic DNA in each sample was confirmed by amplification of a 536 bp fragment of the β-globin gene. Eight of the 60 cell lines contained a reproducible abnormally migrating SSCP band. DNA sequence analysis of clones of these eight abnormally migrating SSCP fragments revealed alteration of the primary sequence in each. One of these eight cell lines, HL-60, had two sites of sequence variation in exon 2 of p16, one of which was a common polymorphism at codon 148 (A148T). This polymorphism, which does not affect p16 function, was also present in the colon carcinoma cell line, KM12. Additional sequence variants not known to be polymorphisms were observed in seven (12%) of the 60 cell lines. HL-60 contained a nonsense mutation at codon 80 and HCT-116 contained a one bp insertion at codon 22-23, which results in a frameshift at codon 22 and termination after codon 42. Both of these mutations were reasoned to cause loss of p16 function. Three cell lines (MDA-

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MB-435, MDA-N, and M14) contained the same splice site mutation [T to C substitution at nucleotide 2 of intron 1 (I1+2^{T-C})], and 2 cell lines (UACC-257 and DU-145) had distinct missense mutations. The splice site mutation resulted in aberrant splicing creating a shortened mRNA that had deletion of codons 28 to 50. The functional effect of the splice site and missense mutations was assessed by measuring the binding of GST-p16 fusion proteins to CDK4. Binding of mutant GST-p16 fusion proteins (I1+2^{T-C}, D84Y, and P81L) to CDK4 was 3.2%, 4.9%, and 34% of the binding ability of normal p16, respectively (p<0.0001 for each comparison, 2-tailed Student t-test). Thus, 36 of 60 (60%) cell lines of the NCI drug screen panel contained a genetic alteration (homozygous deletion or intragenic mutation) of p16 that disrupted the function of p16^{INK4A}.

To detect non-genetic alterations associated with loss of p16 function, p16 mRNA and protein expression were examined. Using RT-PCR and subsequent Southern blot hybridization analyses, p16 mRNA expression was undetectable in 41 of 60 (68%) cell lines examined, including 11 of 24 (46%) without detectable genetic alteration. The amplified p16 cDNAs in two cell lines (MDA-MB-435 and MDA-N) were smaller than expected, consistent with altered mRNA splicing as a result of the I1+2^{T-C} mutation. p16 mRNA was not detected in the third cell line (M14) with this splice site mutation. A protein of 16 kd was detected in 17 of the 60 (28%) cell lines by Western blot analysis using p16 polyclonal antiserum. The cell line with a nonsense mutation (HL60) expressed p16 mRNA but not p16 protein. The two cell lines with missense mutations (UACC-257 and DU-145) expressed both mRNA and protein. In UACC-257, a protein smaller than 16 kd was detected, perhaps the result of altered susceptibility to proteolysis of p16^{P81L}. A protein of 16 kd was detected in two cell lines with the splice site mutation (MDA-MB-435 and MDA-N) but was absent in the third cell line with the $I1+2^{T-C}$ mutation, M14. In each cell line, absent or altered p16 protein could be attributed to mutation or transcriptional suppression. In total, 47 of the 60 (78%) cell lines of the NCI drug screen panel had an alteration of p16.

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Comparison of p16 status with growth inhibitory activity. To identify compounds more active against p16-altered cells than p16-normal cells, the p16 status of the 60 cell lines was matched to the growth inhibitory (GI₅₀) activity of the compounds of the NCI drug screen program and ranked according to Pearson correlation coefficients using the COMPARE algorithm. The growth inhibitory activity of cephalostatin 1, a disteroidal alkaloid extracted from the marine worm, *Cephalodiscus gilchristi*, correlated best with p16 status (r=0.599). The growth inhibitory activity of five related compounds [cephalostatins 7, 9, 8, 4 and 3 were also positively correlated with p16 status (r=0.504, 0.493, 0.491, 0.461, and 0.458, respectively). Bryostatin 1, a protein kinase C activator isolated from the marine bryozoan, *Bugula neritina*, had a correlation coefficient of 0.469.

Aliquots of 26 of the 40 compounds with the highest Pearson correlation rankings were available for further *in vitro* analysis. These compounds were assessed for CDK4:cylin D kinase inhibitory activity using baculovirus-expressed human CDK4 and cyclin D1, and a GST-Rb fusion protein as substrate. Six of the 26 compounds examined inhibited phosphorylation of Rb protein by CDK4:cyclin D1 complex with IC₅₀ values ranging from 6.8 to more than 100 μM. No inhibition of GST-Rb phosphorylation by CDK4:cyclin D1 was observed in the presence of the other 20 compounds at concentrations up to 100 μM. The most potent inhibitor was 3-amino-9-thio(10H)-acridone (3-ATA; Formula 3) with an IC₅₀ of 6.8 μM, a value similar to the mean GI₅₀ (30 μM) observed for this compound in the 2 day growth assay of the NCI drug screen. Cephalostatin 1, which has potent antitum or activity *in vitro* (ED₅₀ 10⁻⁷ to 10⁻⁹ μg/ml), had an IC₅₀ for CDK4:cyclin D1 of 20 μM and bryostatin 1 had no inhibitory activity at the highest concentration examined (100 μM).

Characterization of 3-ATA. To examine the specificity of 3-ATA inhibitory activity for CDK4:cyclin D1 kinase, we performed *in vitro* kinase assays using baculovirus-expressed human CDC2:cyclin A, CDK2:cyclin A, and CDK2:cyclin E complexes. 3-ATA was a less potent inhibitor of CDC2 and CDK2 kinase activities with IC₅₀ values at least nine-fold higher compared to the IC₅₀ for CDK4. The addition of 100 μM 3-ATA decreased the binding of CDK4

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to normal p16 by 70% in the p16-CDK4 binding assay (p<0.0001, 2-tailed Student t-test), suggesting that 3-ATA may be acting by a mechanism similar to p16. In the CDK4 kinase assay, the addition of exogenous ATP (0 to 600 μ M) did not alter the inhibitory activity of 3-ATA, suggesting that 3-ATA was not competing with ATP. Thus, 3-ATA appears to inhibit cyclin-dependent kinase activity by a mechanism distinct from that of the flavone L86827 and butyrolactone I, which are known to compete with ATP.

Identification of CDK4-specific inhibitors. To identify compounds in the NCI drug screen that may have a similar mechanism of action as 3-ATA, the pattern of growth inhibitory activity (GI_{50}) of 3-ATA with the GI_{50} of all previously tested compounds as compared. Six compounds not previously examined for CDK4 kinase inhibitory activity had similar patterns of growth inhibitory activity with correlation coefficients greater than 0.6. Among these six, two benzothiadiazine (BTD) compounds (Compound 6) and NSC 645788) inhibited CDK4:cyclin D1 kinase activity *in vitro* with IC_{50} 's (5.0 and 17 μ M, respectively) similar to the IC_{50} of 3-ATA (6.8 μ M).

An additional 45 compounds with structural similarity to 3-ATA and (Compound 6) were available for analysis. Nineteen of these compounds inhibited CDK4 kinase activity with IC₅₀'s ranging from 1.1 to more than 100 μM. Four compounds, 2 structurally related to 3-ATA (Compound 4) and NSC 645153), and 2, Compound 7 and Compound 8, were more potent CDK4 kinase inhibitors than the parent compounds. Compound 4, Compound 7, and Compound 8 also had no CDC2 or CDK2 kinase inhibitory activity at concentrations up to 100 μM. However, two of these compounds, Compound 4 and Compound 7, did not inhibit p16^{INK4A} binding to CDK4, suggesting that their mechanism of inhibition of CDK4 kinase activity is distinct from 3-ATA.

Example 2

This example describes a method for treating cancer using the compounds of the invention. Thioacridones or benzothiadiazines satisfying Formulas 1 and 2 above are obtained that specifically inhibit CDK4:cyclin kinase such that these compounds have an IC₅₀ for CDK4 that is smaller than their IC₅₀

for CDC2 or CDK2. These compounds are administered intravenously or orally to humans at a dose of between 1 μg and 10 grams, preferable between 1 mg and 900 mg per m² of body surface of the patient. The compounds also can be mixed with at least one additive selected from the group consisting of carriers, diluents, excipients, diagnostics, direct compression buffers, buffers, stabilizers, fillers, disintegrates, flavors, colors, and mixtures thereof to form pharmaceutical compositions. The compositions are administered intravenously or orally to humans at a dose of between 1 μg and 10 grams, preferable between 1 mg and 900 mg per m² of body surface of the patient.

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CELL LINE DATA

Compounds of the present invention have been subjected to the drug screening procedure employed by the National Cancer Institute for the screening of drugs having possible anticancer utility. The screening procedure uses a diverse, disease-oriented panel consisting of different human tumor cell lines organized into disease-specific subpanels. The compounds of the present invention were tested over a range of concentrations for cytotoxic or growth-inhibitory effects against cell lines comprising the panel. The subpanels represented diverse histologies (leukemias, melanomas, and tumors of the lung, colon, kidney, breast, ovary, and brain). The tests produced individual dose-responses, one for each cell line (i.e., one for each example), and the data are disclosed in dose-response curves, e.g., FIGS. 1(A)-1(I). The data provided by these dose response curves are summarized using a mean-graph format, e.g., FIG. 2.

To produce data for the mean-graph format, a compound concentration that produced a target level response was calculated for each cell line. Three different response parameters were evaluated. The first response parameter was the growth inhibition (" GI_{50} "). GI_{50} is the concentration of compounds made according to the present invention that produced an apparent 50% decrease in the number of tumor cells relative to the appropriate control (not exposed to the compounds of the present invention) at the end of the incubation period.

The second response parameter was the total growth inhibition

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("TGI"). TGI is the concentration at which the number of tumor cells remaining at the end of the incubation period substantially equal the number of tumor cells existing at the start of the incubation period.

The third response parameter was the lethal concentration (" LC_{50} "). LC_{50} is the concentration of compounds made according to the present invention that caused an apparent 50 percent reduction in the number of tumor cells relative to the appropriate control (not exposed to the compounds of the present invention) at the start of the incubation period.

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In a typical GI_{50} mean graph the relative position of the vertical reference line along the horizontal concentration axis was obtained by averaging the negative $\log_{10}GI_{50}$ values for all the cell lines tested against the compound. Horizontal bars were then plotted for the individual negative $\log_{10}GI_{50}$ values of each cell line relative to the vertical reference line. The GI_{50} graph thus provides a characteristic fingerprint for the compound, displaying the individual cell lines that are proportionately more sensitive than average (bars extending to the right of the reference line) or proportionately less sensitive than average (bars extending to the left of the reference line). The length of a bar is proportional to the difference between the $\log_{10}GI_{50}$ value obtained with the particular cell line and the mean (represented by the vertical reference line).

The data obtained using the cell line procedures referred to above are provided by FIGS. 1-12. This data shows that the compounds of the present invention inhibit the growth of living cells.

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WE CLAIM:

- 1. 3-amino-9- thio(10H)-acridone.
- An antineoplastic composition comprising a therapeutically
 effective amount of a compound selected from the group consisting of a benzothiadiazine, a thioacridone, or mixtures thereof, the compound having an IC₅₀ for CDK4 of less than about 10 μM.
- 3. The antineoplastic composition of claim 2 wherein the compound has an IC₅₀ for CDC2 of more than about 60 μ M.
 - 4. The antineoplastic composition of claim 3 wherein the compound has an IC₅₀ for CDK2/A of more than about 100 μ M.
- 5. The antineoplastic composition of claim 4 wherein the compound has an IC₅₀ for CDK2/E of more than about 80 μ M.
 - 6. The antineoplastic composition of claim 2 wherein the compound has an IC $_{50}$ for CDK4 of less than about 2.5 μM .

7. The antineoplastic composition of claim 3 wherein the compound further has an IC₅₀ for CDC2 of greater than about 100 μ M.

- 8. The antineoplastic composition of claim 7 wherein the compound further has an IC_{50} for CDK2/A of greater than about 100 μ M.
 - 9. The antineoplastic composition of claim 8 wherein the compound further has an IC₅₀ for CDK2/E of greater than about 100 μ M.
- 30 10. An antineoplastic composition comprising a compound selected from the group consisting of a benzothiadiazine, a thioacridone, or mixtures thereof, the compound having an IC₅₀ ratio for CDC2:CDK4 of greater than

about 8.5.

11. The antineoplastic composition of claim 10, wherein the compound has an IC₅₀ ratio for CDK2/A:CDK4 of greater than about 14.

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- 12. The antineoplastic composition of claim 11, wherein the compound has an IC₅₀ ratio for CDC2/E:CDK4 of greater than about 11.5.
- 13. The antineoplastic composition of claim 10 wherein the compound has an IC₅₀ ratio for CDC2:CDK4 of greater than about 20, an IC₅₀ ratio for CDK2/A:CDK4 of greater than about 20, and an IC₅₀ ratio for CDC2/E:CDK4 of greater than about 20.
- 14. The antineoplastic composition of claim 10 wherein the compound
 15 has an IC₅₀ ratio for CDC2:CDK4 of greater than about 60, an IC₅₀ ratio for CDK2/A:CDK4 of greater than about 60, and an IC₅₀ ratio for CDC2/E:CDK4 of greater than about 60.
- 15. An antineoplastic composition comprising an effective amount of a compound according to Formula 1

where m is 0 or 1, n = m, R_1 - R_4 are independently selected from the group consisting of H, -NH₂ and lower alkoxy, where with m=1 one of R_1 - R_4 is an amine bonded to R= to form an arylamide,

Formula 2

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where R and R_1 are independently carbon or nitrogen, where if R_1 =carbon X is hydrogen or halogen, and R_2 is selected from the group consisting of lower alkyl and aryl amino,

and mixtures of compounds satisfying Formula 1 and/or Formula 2, wherein the compounds have an IC₅₀ for CDK4 of less than about 10 μ M.

- 15 16. The composition according to claim 15 wherein the compound has an IC₅₀ for CDC2 of more than about 60 μ M.
 - 17. The composition according to claim 15 wherein the compound has an IC $_{50}$ for CDK2/A of greater than about 100 μ M.

- 18. The composition according to claim 15 wherein the compound has an IC $_{50}$ for CDK2/E of greater than about 80 μM .
- 19. The composition according to claim 15 and further comprising additives selected from the group consisting of carriers, diluents, excipients, diagnostics, direct compression buffers, buffers, stabilizers, fillers, disintegrates, flavors, colors, and mixtures thereof.
- 20. The composition according to claim 15 wherein the effective amount of the compound is sufficient to provide from about 1 mg to about 900 mg per m² body surface area of a subject treated with the composition.

21. The composition according to claim 13 comprising

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22. The composition according to claim 15 where, with respect to Formula 1, m=n=0.

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- 23. The composition according to claim 22 where at least one of R_1 - R_4 is an amine, the remainder of R_1 - R_4 being hydrogen.
- 24. The composition according to claim 22 comprising 3-amino-9-thio(10H)-acridone.
 - 25. The composition according to claim 22 where at least one of R_1 - R_4 is lower alkoxy.
- 25 26. The composition according to claim 22 where at least two of R_1 - R_4 are lower alkoxy.
 - 27. The composition according to claim 22 where at least two of R_1 - R_4 are methoxy.

28. The composition according to claim 22 comprising

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- 29. The composition according to claim 15 where, with respect to 10 Formula 2, R is nitrogen.
 - 30. The composition according to claim 27 where R_2 is a lower alkyl.
- 31. The composition according to claim 28 where R_2 is selected from the group consisting of methyl and ethyl.
 - 32. The composition according to claim 27 where X is halogen.
 - 33. The composition according to claim 28 where X is halogen.

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34. The composition according to claim 31 comprising

35. The composition according to claim 31 comprising

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- 36. The composition according to claim 15 where the compound has an IC₅₀ for CDK4/D1 of less than about 10 μ M.
 - 37. The composition according to claim 15 where the compound has an IC₅₀ for CDK4/D1 of from about 1 μ M to about 7 μ M.
- 15 38. The composition according to claim 35 where the compound has IC_{50} values for CDC2/A and CDK2/A of greater than about 60 μ M.
 - 39. A method for inhibiting the growth of living cells, comprising: providing a compound selected from the group consisting of a benzothiadiazine, a thioacridone, or mixtures thereof, the compound having an IC₅₀ for CDK4 of less than about 10 μM, an IC₅₀ for CDC2 of more than about 60 μM, an IC₅₀ for CDK2/A of more than about 100 μM, and an IC₅₀ for CDK2/E of more than about 80 μM; and

administering an effective amount of the compound to inhibit the growth of living cells.

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40. A method for inhibiting the growth of living cells, comprising: providing a compound selected from the group consisting of a benzothiadiazine, a thioacridone, or mixtures thereof, the compound having the compound having an IC₅₀ ratio for CDC2:CDK4 of greater than about 8.5, an IC₅₀ ratio for CDK2/A:CDK4 of greater than about 14, and an IC₅₀ ratio for CDC2/E:CDK4 of greater than about 11.5; and

administering an effective amount of the compound to inhibit the growth of living cells.

10 41. The method according to claim 40 where the compound satisfies Formula 1

where m is 0 or 1, n = m, R_1 - R_4 are independently selected from the group consisting of H, -NH₂ and lower alkoxy, where with m=1 R= is an arylamide bonded to one of R_1 - R_4 which is an amine, or Formula 2

where R and R_1 are independently carbon or nitrogen, where if R_1 =carbon X is hydrogen or halogen, and R_2 is selected from the group consisting of lower alkyl

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and aryl amino, and mixtures of compounds satisfying Formula 1 and/or Formula 2.

- 42. The method according to claim 39 where the step of providing a compound comprises providing a composition comprising the compound, and further comprising additives selected from the group consisting of carriers, diluents, excipients, diagnostics, direct compression buffers, buffers, stabilizers, fillers, disintegrates, flavors, colors, and mixtures thereof.
 - 43. The method according to claim 41 comprising

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- 44. The method according to claim 41 where, with respect to Formula 1, m=n=0.
- 25 45. The method according to claim 44 where at least one of R_1 - R_4 is an amine, the remainder of R_1 - R_4 being hydrogen.
 - 46. The method according to claim 44 comprising 3-amino-9-thio(10H)-acridone.

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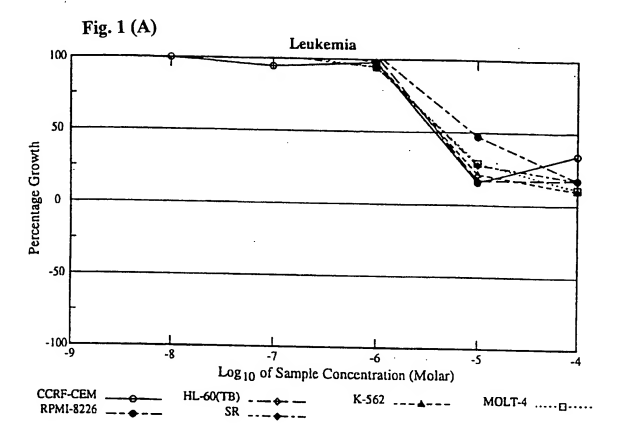
47. The method according to claim 41 where at least one of R_1 - R_4 is lower alkoxy.

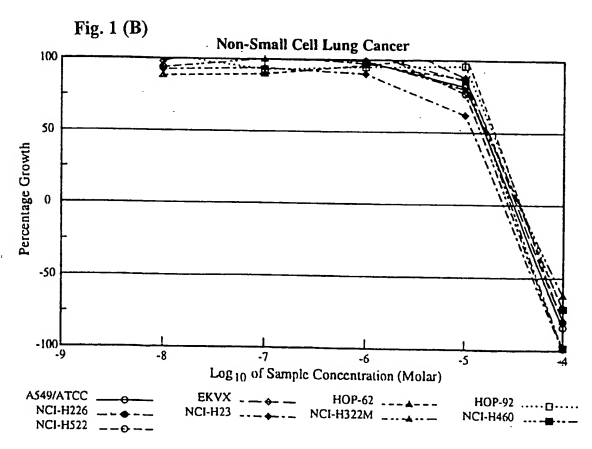
- 48. The method according to claim 41 where at least two of R_1 - R_4 are lower alkoxy.
- 49. The method according to claim 41 where at least two of R_1 - R_4 are methoxy.
 - 50. The method according to claim 49 comprising

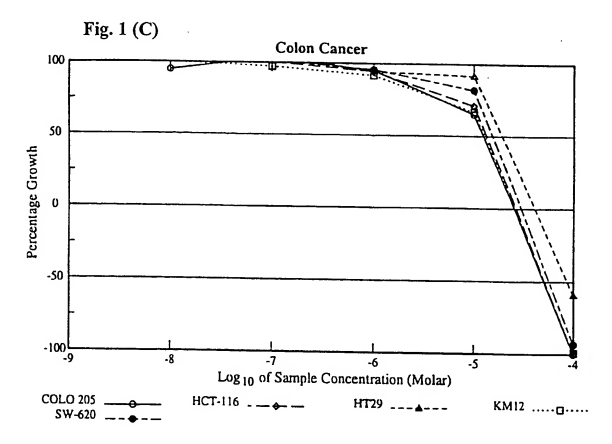
- 15 51. The method according to claim 41 where, with respect to Formula 2, R is nitrogen.
 - 52. The method according to claim 51 where R2 is lower alkyl.
- 53. The method according to claim 52 where R_2 is selected from the group consisting of methyl and ethyl.
 - 54. The method according to claim 51 where X is halogen.
- 25 55. The method according to claim 52 where X is halogen.

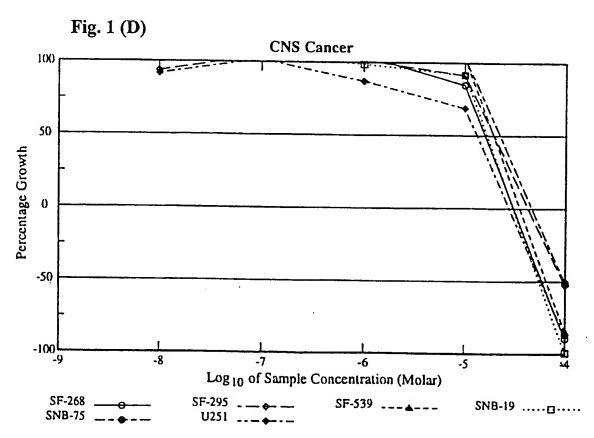
56. The method according to claim 51 comprising

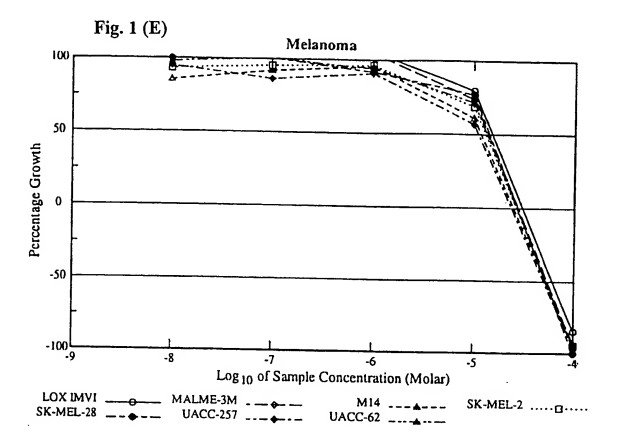
57. The method according to claim 51 comprising

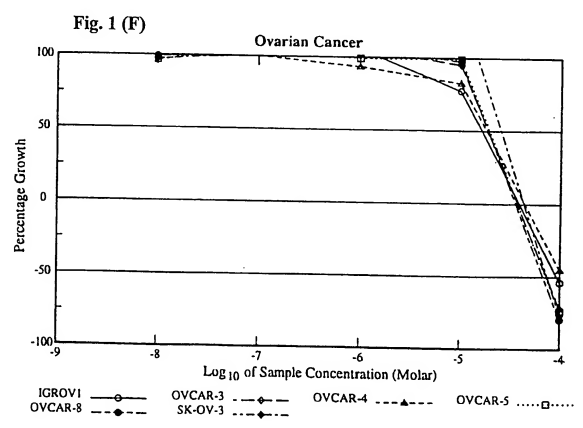


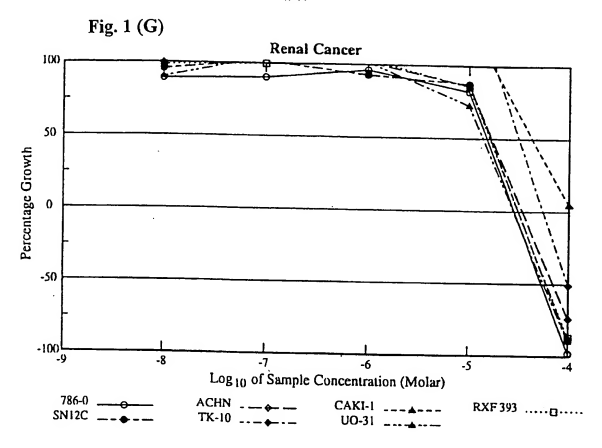


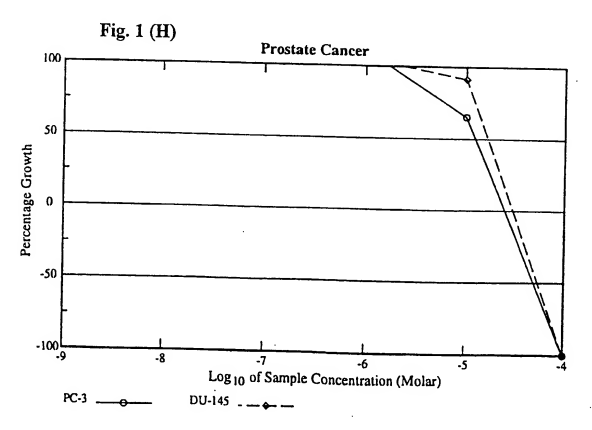












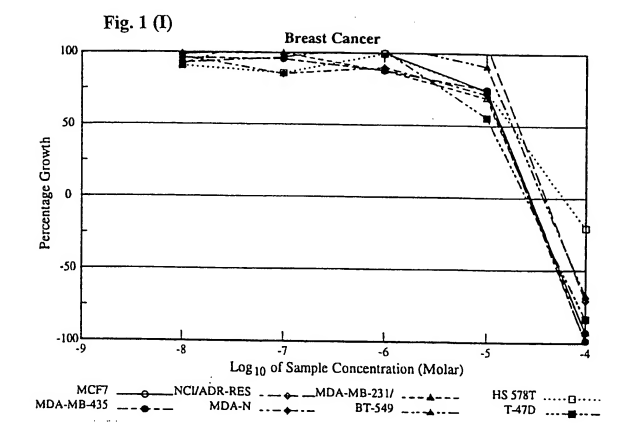


Fig. 2 (Sheet 1 of 3)

Panel/Cell Line	Log ₁₀ GI50	G150
Leukemia		1
CCRF-CEM	-5.42	
HL-60(TB)	-5.39	
K-562	-5.37	
MOLT-4	-5.32	
RPMI-8226	-5.05	
SR .	-5.33	
Non-Small Cell Lung Cancer	***************************************	•••••••••••••••••••••••••••••••••••••••
A549/ATCC	-4.81	•
EKVX	-4.77	
HOP-62	-4.71	•
HOP-92	-4.73	
NCI-H226	-4.77	•
NCI-H23	-4.92	
NCI-H322M	-4.79	
NCI-H460	-4.81	
NCI-H522	-4.85	1
Colon Cancer	••••••••••••	
COLO 205	-4.91	•
HCT-116	-4.88)
HT29	-4.72	
KM12	-4.90	•
SW-620	-4.82	
CNS Cancer		
SF-268	-4.80	
SF-295	-4.71	
SF-539	-4.72	
SNB-19	-4.78	
SNB-75	-4.65	2
U251	-4.88	
Melanoma	-4.60	
LOX IMVI	-4.82	
MALME-3M	-4.86	4
M14	-4.93	
SK-MEL-2	-4.89	
SK-MEL-28	•	í
UACC-257	-4.85	Ļ
UACC-62	-4.95	Γ
Ovarian Cancer	-4.87	
IGROVI	-4.79	
OVCAR-3		
OVCAR-4	-4.73	
OVCAR-5	-4.74	
	-4.72	
OVCAR-8	-4.73	
SK-OV-3	-4.60	0.00
Renal Cancer		
786-0	-4.82	1
ACHN	-4.77	
CAKI-I	-4.36	
RXF 393	-4.79]
SN12C	-4.79	
TK-10	-4.50	
UO-31	-4.86	
Prostate Cancer		
PC-3	-4.91	<u>}</u>
DU-145	-4.78	1
Breast Cancer		
MCF7	-4.86	
NCVADR-RES	-4.68	4
MDA-MB-231/ATCC	-4.88	1
HS 578T	-4.75	•
MDA-MB-435	-4.86	
MDA-N	-4.87	1
BT-549	-4.74	•
T-47D	-4.96	þ
MG_MID	-4.85	· L
Delta	0.57	.
Range	1.05	
	1 1	1 1 1
	+3 +2	2 +1 0 -1 -2 -3

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Fig. 2 (Sheet 2 of 3)

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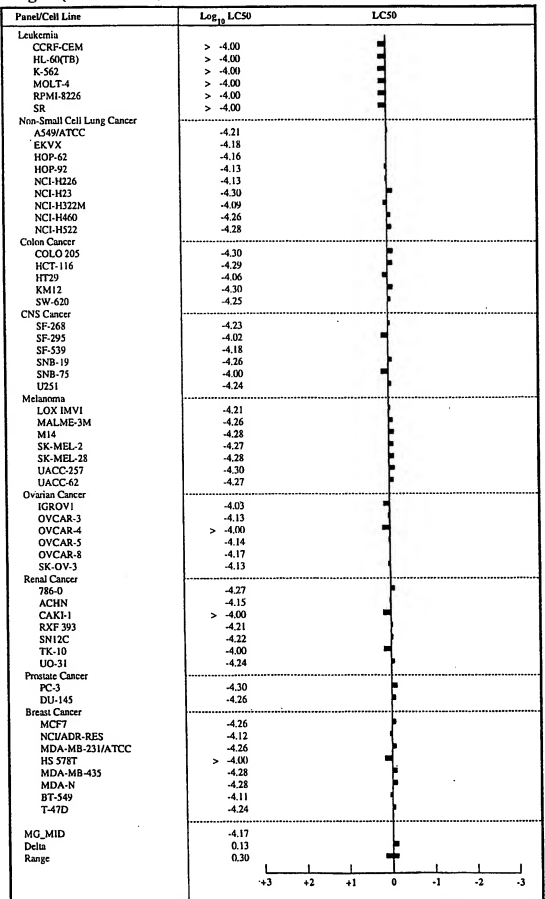
Panel/Cell Line	Log ₁₀ TGI	TGI	
Leukemia			
CCRF-CEM	> -4.00		
HL-60(TB)	> -4.00		
K-562	> -4.00	T-STATE OF THE STATE OF THE STA	
MOLT-4	> -4.00		
RPMI-8226	> -4.00		
SR	> -4.00		
Non-Small Cell Lung Cancer	***************************************		
A549/ATCC	-4.51		
EKVX	-4.47	Ţ.	
HOP-62	-4.43	ì	
HOP-92	-4.43		
NCI-H226	-4.45 -4.45		
NCI-H23	-4.43 -4.61		
NCI-H322M	•		
NCI-H460	-4.44	L	
NCI-H400 NCI-H522	-4.53		
	-4.56		
Colon Cancer	***************************************	***************************************	
COLO 205	-4.61		
HCT-116	-4.59		
НТ29	-4.39		
KM12	-4.60		
SW-620	-4.53		
CNS Cancer			
SF-268	-4,51		.~~~~
SF-295	-4.37		
SF-539	-4.45		
SNB-19	-4.52		
SNB-75	-4.33	<u> </u>	
U251			
Melanoma	-4.56		
LOX IMVI	4.50		
	-4.52		
MALME-3M M14	-4.56	Ja Pa	
	-4.61		
SK-MEL-2	-4.58		
SK-MEL-28	-4.57	•	
UACC-257	-4.63		
UACC-62	-4.57		
Ovarian Cancer	***************************************		
IGROVI	-4.41		
OVCAR-3	-4.43	0.00	
OVCAR-4	-4.35	•	
OVCAR-5	-4.43		
OVCAR-8	-4.45	ì	
SK-OV-3	-4.36	4	
Renal Cancer	*****************************	***************************************	
786-0	-4.54		
ACHN	-4.46		
CAKI-I	> -4.00		
RXF 393	-4.50		
SN12C	-4.50		
TK-10	-4.25		
UO-31	-4.25 -4.55		
Prostate Cancer			
PC-3	4.41		•••
DU-145	-4.61		
Breast Cancer	-4.52		
			•••••
MCF7	-4.56		
NCVADR-RES	-4.40	9 7 10 20	
MDA-MB-231/ATCC	-4.57	- N. C.	
HS 578T	-4.22		
MDA-MB-435	-4.57	V	
MDA-N	-4.58		
BT-549	-4.43		
T-47D	-4.60		

MG_MID	-4.43		
Delta	0.19	_	
Range	0.63		
o-	1 0.03	<u> </u>	
	+3 +2	+1 0 -1 -2	ļ

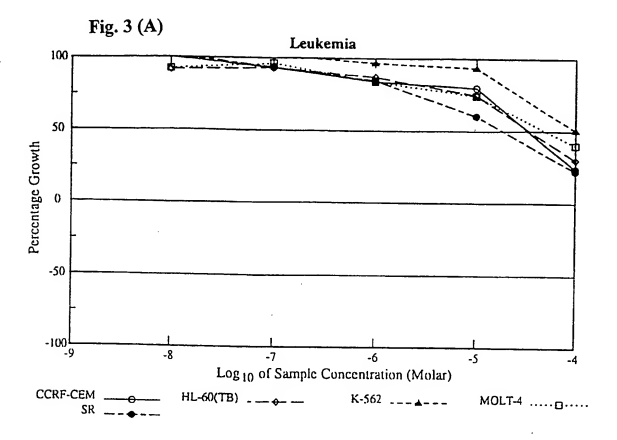
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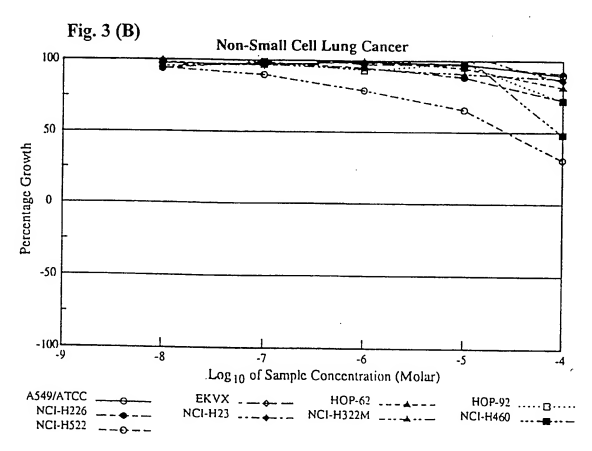
Fig. 2 (Sheet 3 of 3)

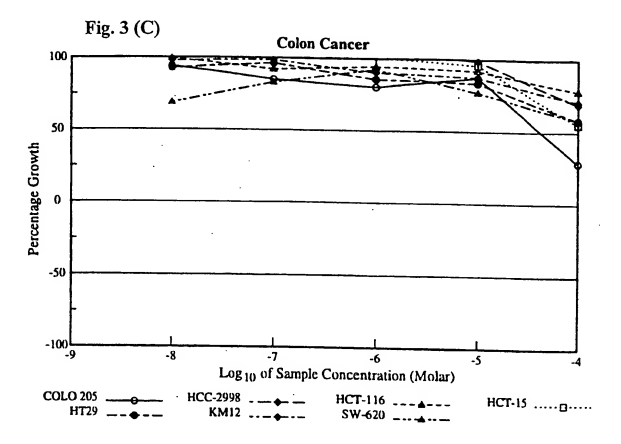
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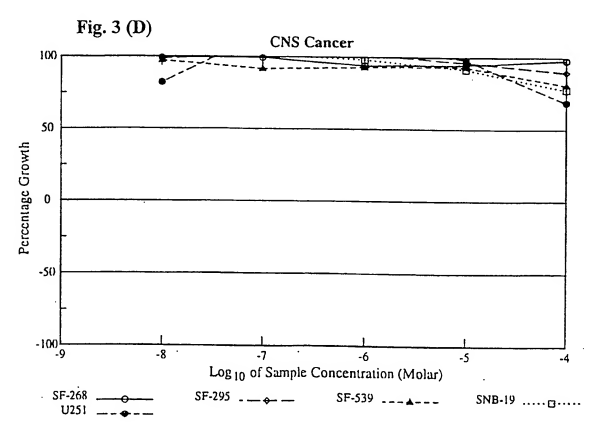


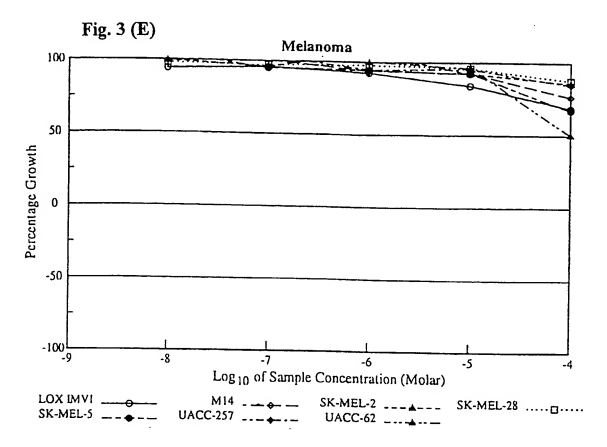
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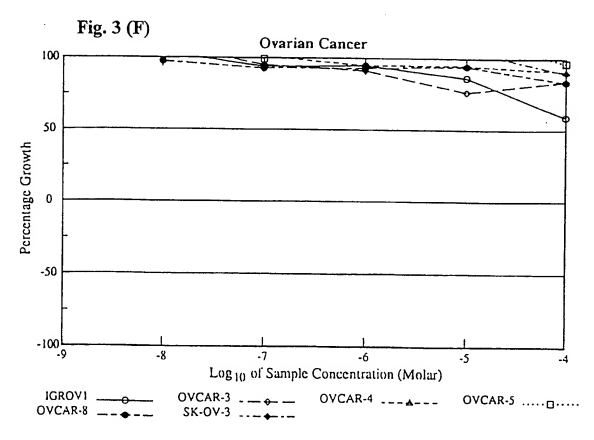


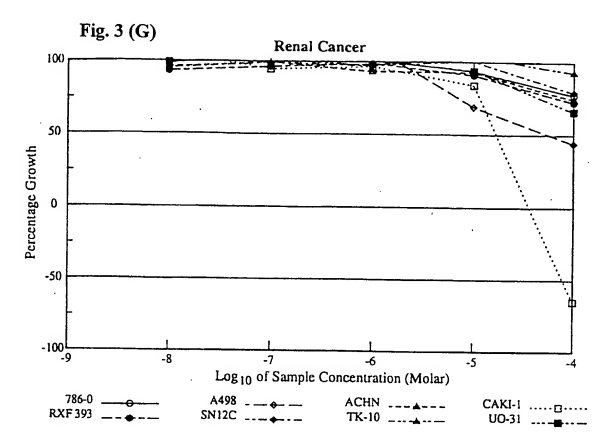


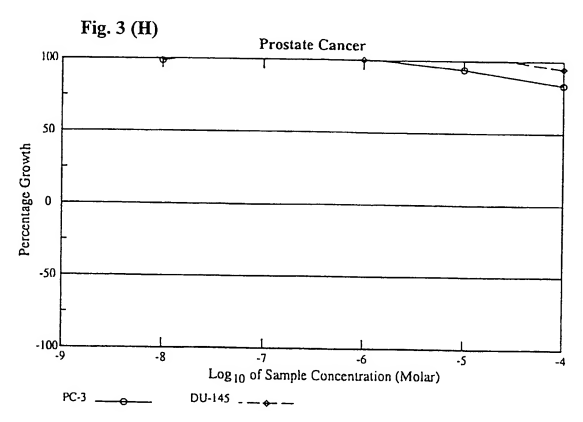












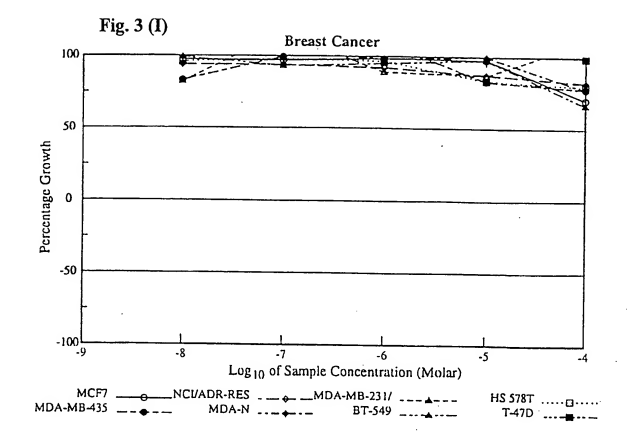


Fig. 4 (Sheet 1 of 3) 14/47

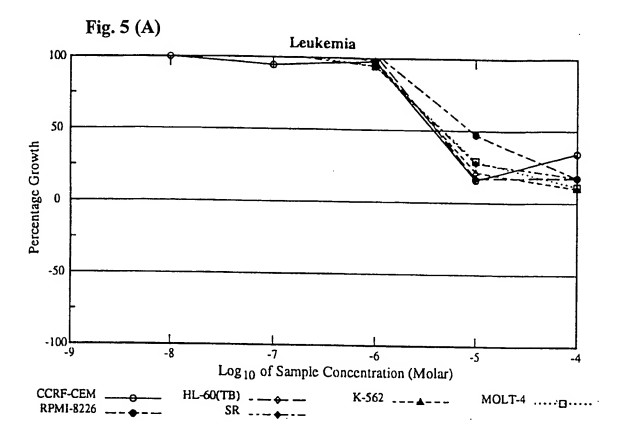
Panel/Cell Line Los ₁₀₀ Gi50 Gi50	Fig. 4 (Sneet 1 of 3)	17	/4/			
Leukemia CCRF-CEM H-60(TB) A-45 A-	Panel/Cell Line	Logio G150)	G	150	
H.L-60(TB)	Leukemia					
H60(TB)	CCRF-CEM	-4.47				
MOLT-4 SR Non-Small Cell Lung Cancer ASAMATCC EXVX	HL-60(TB)	· ·			-	
MOLT-4 SR Non-Small Cell Lung Cancer A549/ATCC EKVX					4	
SR 4.73 Non-Small Cell Lung Cancer A549/ATCC	MOLT-4	· ·			-	
Non-Small Cell Lung Cancer A549ATCC EXVX	SR					
ASMATCC ERVX			•••••••			• • • • • • • • • • • • • • • • • • • •
EKYX		> -4 (x)				
HOP-92 HOP-92 NCI-H236 NCI-H231 NCI-H2460 NCI-H252 NCI-H254 NCI-H360 NCI-H3					1	
HOP-92 NCI-H226 NCI-H236 NCI-H321 NCI-H321 NCI-H321 NCI-H340 NCI-H361 NCI-H361 NCI-H362 NCI-H362 NCI-H362 NCI-H364 NCI-H362 NCI-H364 NCI-H362 NCI-H364 NCI-H362 NCI-H364 NCI-H362 NCI-H364 NCI-H]	
NCI-H23 NCI-H23 NCI-H240 NCI-H240 NCI-H321 NCI-H360 NCI-H322 NCI-H360 NCI-H522 -4.03 NCI-H522 -4.35 Colon Canter COLO 205 HCT-116 > 4.00 HCT-116 > 4.00 HCT-15 > 4.00 HCT-15 > 4.00 HCT-179 > 4.00 SW-620 CNS Cancer SF-268 SF-395 SF-395 SR-19 U251 Nd-hanoma LOX IMVI NIA LOX IMVI NIA SK-MEL-2 S]	
NCI-H22M NCI-H32DM NCI-H360 NCI-H360 NCI-H361 NCI-H362 NCI-H362 NCI-H360 NCI-H362 NC						
NCI-H322M NCI-H460 NCI-H522 NCI-H6460 NCI-H522 A.55 Colon Cancer COLO 205 HCC.2998 A.4.00 HCT-116 A.4.00 HCT-15 A.4.00 HCT-15 A.4.00 HCT-15 A.4.00 HCT-17 KM12 A.4.00 SW-620 CNS Cancer SF-268 SF-295 SF-268 SF-295 SR-399 SNB-19 A.00 SS-339 SNB-19 A.00 SS-339 SNB-19 A.00 McI-Androma LOX IMVI M14 SK-MEL-2 SK-MEL-3 SK-MEL						
NCI-H460 NCI-H522 Colon Canner COLO 205 HCC-2998 HCC-2998 HCC-1998 HCC-115 HCC-2998 HCC-115 HCC-2998 HCC-115 HCC-2998 HCC-115 HCC-2998 HCC-115 HCC-2998 HCC-115 HCC-116 HCC-115 HCC-116 HCC-115 HCC-116 HCC-115 HCC-116 HCC-115 HCC-116 HCC-115 HCC-116 HCC-11]	
NCH-B52 Colon Cancer COLO 205 HCC-2998 HCT-116 S					1	
Color Cancer COLO 205 HCC-3998 HCT-116 HCT-15 HCT-1					<u> </u>	
COLO 2015 HCC-3998 HCT-116 HCT-15 HCC-3998 HCT-116 HCT-15 HT29 Au0 HT29 XM-20		-4.33				
HCC-3998 HCT-116 HCT-15 HCT-16 HCT-15 HCT-16 HCT-15 HCT-16 HCT-15 HCT-16 HCT-15 HCT-16		4.00	•••••••	••••••	·	
HCT-116 HCT-15 HCT-15 HCT-15 HCT-19 HCT-15 HCT-19 H					_	
HCT-15 HT29 KM12 HT39 KM12 SW-620 SW-620 SW-620 SW-620 SF-268 SF-268 SF-295 SF-399 SNB-19 U251 A-00 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-23 SK-MEL-23 SK-MEL-25 UACC-62 UACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-63 VARACC-64 OVCAR-3 OVCAR-4 OVCAR-3 OVCAR-4 OVCAR-5 SK-MCL-3 SK-MCL-3 SK-MCL-3 SK-MCL-3 VARACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-62 VARACC-63 VARACC-64 VARACC-64 VARACC-65 VARACC-65 VARACC-66 VARACCC-66 VARACCC-67 VARACCC-68 VARACCCC-68 VARACCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC		-			1	
HTT9					3	
KM12					1	
SW-620 CNS Cancer SF-268 SF-295 SF-399 SNB-19 U251 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-3 SK-MEL-5 UACC-62 VACC-62 VACC-62 VACC-62 VACC-63 VACC-64 OVCAR-4 OVCAR-3 OVCAR-4 OVCAR-5 SK-OV-3 SK-OV-3 Renal Cancer T86-0 A498 ACHN CAKI-1 CAMICA CA					1	
CNS Cancer SF-268 SF-295 SF-399 SNB-19 U251 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-2 SK-MEL-28 SK-MEL-3 UACC-257 UACC-262 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 SK-OV-3 SK-OV-3 SK-OV-3 SK-OV-3 SR-01 A498 A498 ACHN CAKL-1 4.77 RXF 393 SNIZC TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS-400 MDA-MB-231/ATCC HS-578T H-000 MG_MID Delta Range MG_MID Delta Range MG_MID Delta Range A00 A407 A					1	
SF-268 SF-295 SF-399 SNB-19 U251 V251 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-3 SK-MEL-3 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 SK-OV-3 Renal Cancer 786-0 A498 A-423 ACHN CAKI-1 RXF 393 SN12C TK-10 UO-31 Prostate Cancer MCF7 NCVADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-MB-435 MDA-MB-435 MDA-MB-435 MC-MID Delta MG_MID Delta Range MG_MID Delta Range MG_MID Delta Range - 4.00 - 4.00 -		> -4.00			1	
SF-295 SF-339 SNB-19 U251 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-2 SK-MEL-28 SK-MEL-5 UACC-257 UACC-257 UACC-262 VOYAR-4 OVCAR-3 OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 RXF 393 SN12C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 MCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 78-99 MDA-MB-2		• • • • • • • • • • • • • • • • • • • •	••••••••••••••			*************
SF.295 SF.399 SNB-19 U251 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-28 SK-MEL-28 SK-MEL-3 > 4.00 UACC-257 UACC-62 UACC-262 UACC-27 UACC-62 UACC-42 OVGAR-3 OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-5 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 RXF 393 SNI2C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 MCL/ADR-RES MDA-MB-231/ATCC HS 578T N-4,00 MG_MID Delta DO-10 MG_MID Delta Range MC_MID MG_MID Delta Range MCMMID Delta Range MCMMID Delta Range -4.00 -4.00 -4.0		> -4.00			4	
SF-539 SNB-19 U251 Nelanoma LOX IMVI M14 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-5 UACC-257 UACC-257 UACC-26 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-5 SK-OV-3 SK-OV-3 Renal Cancer 786-0 A498 A498 A498 A498 ACHN CAKI-1 RF-393 SNI2C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-MB-435 MDA-MB-435 MG_MID Delta MG_MID Delta MG_MID Delta MG_MID Delta Range MG_MID Delta Range MG_MID Delta Range MG_MID Delta Range A-00 A-0	SF-295	> -4.00			₹	
SNB-19	SF-539				4	
U251	SNB-19				d	
Melanoma LOX IMVI	U251				4	
LOX IMVI M14 SK-MEL-2 SK-MEL-28 SK-MEL-5 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 -4.00 CAKI-1 RXF 393 SNI2C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-23I/ATCC HS 578T MDA-MB-435 MDA-M MDA-MB MDA-M MDA-MB MDA-			•			
MI4 SK-MEL-2 SK-MEL-2 SK-MEL-5 SK-MEL-5 SK-MEL-5 UACC-257 UACC-27 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-5 OVCAR-8 SK-OV-3 SK-OV-3 Renal Cancer 786-0 A4.00 CAKI-1 R-KF 393 ACHN CAKI-1 R-KF 393 SN-12C TK-10 U-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC MDA-MB-35 MDA-MB		> 400				
SK-MEL-2 SK-MEL-5 SK-MEL-5 VA.00 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 SK-OV-3 Renal Cancer 786-0 A498 A-400 CAKI-1 RXF 393 SNI2C TK-10 UO-31 VO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-23I/ATCC HS 578T MDA-MB-435 MDA-MB-435 MDA-MB-435 MDA-MB-455 MDA-MB BT-549 T-47D MG_MID Delta Range MG_MID Delta Range - 4.00 - 4.00 - 4.00 - 4.00 - 4.00 - 4.00 - 4.00 - 4.00					1	
SK-MEL-28 SK-MEL-5 SK-MEL-5 VACC-257 VACC-62 VACC-62 VACC-62 VACC-62 VACC-62 VACC-62 VACC-62 VACC-62 VACC-62 VACCC-62 VACCC-62 VACCC-62 VACCC-62 VACCC-62 VACCC VACCC-62 VACCC]	
SK-MEL-5 UACC-257 UACC-27 VACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 SK-OV-3 Renal Cancer 786-0 A498 A23 ACHN CAKI-1 RYF 393 SN12C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 5787 MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta Range MG_MID Delta Range V -4.00 V-4.00 V-4.00 V-4.00 V-4.00 V-4.00 V-6.00 V-6.]	
UACC-257 UACC-62 Varian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 A498 ACHN CAKI-1 CAKI-1 CU-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta MG_MID Delta RA00 VAR-0 VA.00 VAR-0 V]	
UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 RXF 393 SN12C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 578T MDA-MB-231/ATCC HS 578T MDA-MB-235 MDA-N BT-549 T-47D MG_MID Delta Range MGMID Delta Range - 4.00 - 4.00 - 4.00					1	
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OVCAR-5 > 4.00 OVCAR-8 > 4.00 SK-OV-3 > 4.00 Renal Cancer					1	
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786-0 A498 A498 ACHN CAKI-1 CAKI-1 RXF 393 SN12C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta Range > 4.00 A4.00 -4.23 A-4.00 -4.77 A-4.00 -		> -4.00			4	
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TK-10 UO-31 VO-31 Prostate Cancer PC-3 DU-145 Seast Cancer MCF7 NCVADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N ST-549 T-47D MG_MID Delta Range > 4,00 > 4,00 > 4,00 > 4,00		> -4.00			•	
UO-31 > -4.00 Prostate Cancer > -4.00 DU-145 > -4.00 Breast Cancer > -4.00 MCF7 > -4.00 NCVADR-RES > -4.00 MDA-MB-23I/ATCC > -4.00 HS 578T > -4.00 MDA-MB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.07 Delta 0.70 Range 0.77		> -4.00			•	
D0-31		> -4.00			•	
Prostate Cancer PC-3 DU-145 Sreast Cancer MCF7 NCVADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta Range > -4.00 > -4.00 > -4.00					4	
DU-145 > 4,00 Breast Cancer > 4,00 MCF7 > 4,00 NCI/ADR-RES > 4,00 MDA-MB-231/ATCC > 4,00 HS 578T > 4,00 MDA-MB-435 > 4,00 MDA-N > 4,00 BT-549 > 4,00 T-47D > 4,00 MG_MID 4,07 Delta 0.70 Range 0.77				••••••		***************************************
DU-145 > 4.00 Breast Cancer > 4.00 MCF7 > 4.00 MCWADR-RES > 4.00 MDA-MB-231/ATCC > 4.00 HS 578T > 4.00 MDA-MB-435 > 4.00 MDA-N > 4.00 BT-549 > 4.00 T-47D > 4.00 MG_MID 4.07 Delta 0.70 Range 0.77	PC-3	> -4.00			4	
Breast Cancer A.00 MCF7 > 4.00 NCVADR-RES > 4.00 MDA-MB-231/ATCC > 4.00 HS 578T > 4.00 MDA-MB-435 > 4.00 MDA-N > 4.00 BT-549 > 4.00 T-47D > 4.00 MG_MID 4.07 Delta 0.70 Range 0.77	DU-145				4	
MCF7 > 4.00 NCVADR-RES > 4.00 MDA-MB-231/ATCC > 4.00 HS 578T > 4.00 MDA-MB-435 > 4.00 MDA-N > 4.00 BT-549 > 4.00 T-47D > 4.00 MG_MID 4.07 Delta 0.70 Range 0.77	Breast Cancer			*********		******
NCI/ADR-RES > 4,00 MDA-MB-231/ATCC > 4,00 HS 578T > 4,00 MDA-MB-435 > 4,00 MDA-N > 4,00 BT-549 > 4,00 T-47D > 4,00 MG_MID -4,07 Delta 0,70 Range 0,77	MCF7	> 400			4	
MDA-MB-231/ATCC HS 578T		1			4	
HS 578T		i e			į.	
MDA-MB-435		l .			1	
MDA-N BT-549 T-47D MG_MID Delta Range MDA-N > -4.00 > -4.00 -4.07 0.70					1	
BT-549 T-47D MG_MID Delta Range DOI: 10.77 DOI: 10.		1			1	
T-47D > -4.00 MG_MID		1]	
MG_MID -4.07 Delta 0.70 Range 0.77		E .]	
Delta 0.70 Range 0.77	1~4/13	> -4.00			1	
Delta 0.70 Range 0.77	MC MID			*************		
Range 0.77					·	
				·		
+3 +2 +1 0 -1 -2	Kange	0.77				
+3 +2 +1 0 -1 -2		}	<u> </u>			
		1	+3 +2	+1	0 -1	-2 -3

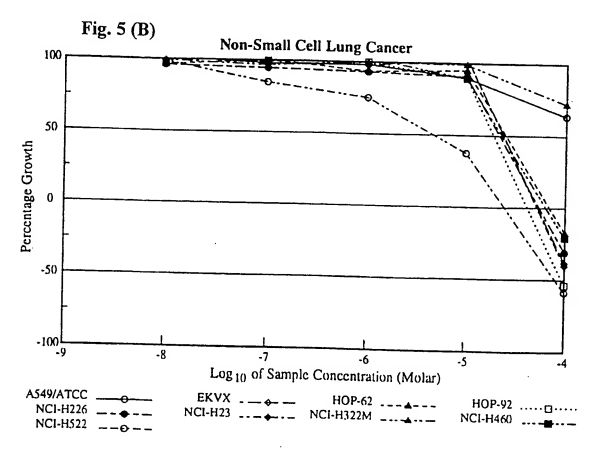
Fig. 4 (Sheet 2 of 3)

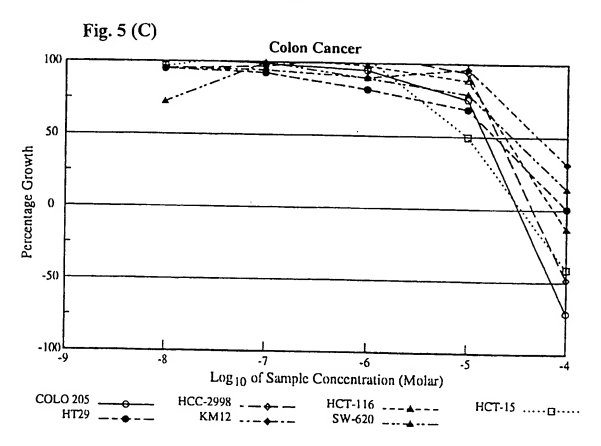
Panel/Cell Line	Log ₁₀ TGI	TGI
Leukemia -		1
CCRF-CEM	> -4.00	
HL-60(TB)	> -4.00	· ·
K-562	> -4.00	
MOLT-4	> -4.00	
SR	> -4.00	
Non-Small Cell Lung Cancer		***************************************
A549/ATCC	> -4.00	
EKVX	> -4.00	
HOP-62	> -4.00	
HOP-92	> -4.00	
NCI-H226	> -4.00	
NCI-H23	> -4.00	
NCI-H322M	> -4.00	
NCI-11460	> -4.00	i
NCI-H522	> -4.00	
Colon Cancer	Z 4,00	
COLO 205	> -4.00	'
HCC-2998	> -4.00	
HCT-116	> 4.00	
HCT-15	> -4.00	
HT29	> -4.00	
KM12	> -4.00	
SW-620		
CNS Cancer	> -4.00	
	4 00	
SF-268	> -4.00	
SF-295	> -4.00	•
SF-539	> -4.00	
SNB-19	> -4.00	
U251	> -4.00	i i i i i i i i i i i i i i i i i i i
Melanoma		•••••••••••••••••••••••••••••••••••••••
LOX IMVI	> -4.00	
M14	> -4.00	
SK-MEL-2	> -4.00	
SK-MEL-28	> -4.00	
SK-MEL-5	> -4.00	
UACC-257	> -4.00	
UACC-62	> -4.00	
Ovarian Cancer	***************************************	•••••••••••••••••••••••••••••••••••••••
IGROVI	> -4.00	
OVCAR-3	> -4.00	
OVCAR-4	> -4.00	
OVCAR-5	> -4.00	
OVCAR-8	> -4.00	
SK-OV-3	> -4.00	
Renal Cancer	***************************************	***************************************
786-0	> -4.00	
A498	> -4.00	1
ACHN	> -4.00	j
CAKI-I	-4.44	—
RXF 393	> -4.00	
SN12C	> -4.00	
TK-10	> -4.00	
UO-31	> -4.00	[
Prostate Cancer		
PC-3	> -4.00	1
DU-145	> -4.00	[
Breast Cancer		
MCF7	> -4.00	· 1
NCVADR-RES	> -4.00	
MDA-MB-231/ATCC	> -4.00	
HS 578T	> -4.00	1
MDA-MB-435	> -4.00	1
MDA-N	> 4.00	1
MDA-N BT-549	> -4.00	}
T-47D	> -4.00	1
1-4/10	> -4.00	
MC MID	-4.01	
MG_MID Delta	0.43	
	0.44	
Range	J V.44	
•	+3	2 +1 0 -1 -2 -3
	1 73 4	r4 TI () al .7 a3

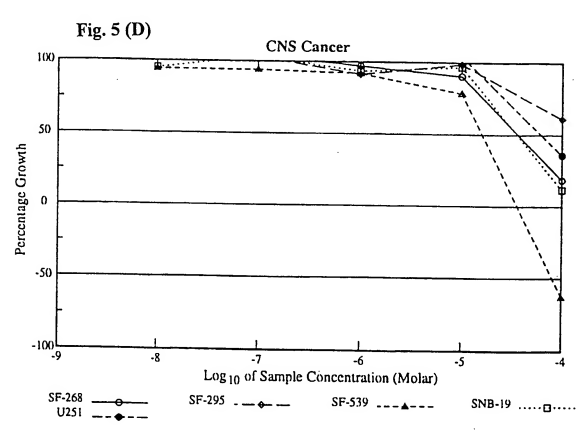
Fig. 4 (Sheet 3 of 3)

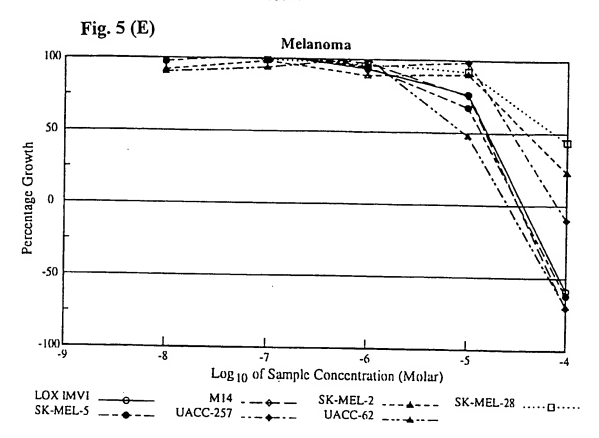
Panel/Cell Line	Log ₁₀ LC50	LC	C50
Leukemia			1
CCRF-CEM	> -4.00		l .
HL-60(TB)	> -4.00		1
K-562	> -4.00		·
MOLT-4	> -4.00		
SR	> -4.00		
Non-Small Cell Lung Cancer	***************************************		***************************************
AS49/ATCC	> -4.00		
EKVX	> -4.00		
HOP-62	> -4.00		
HOP-92 NCI-H226	> -4.00 > -4.00		
NCI-H23	> -4.00		į
NCI-H322M	> -4.00		
NCI-H460	> -4.00		
NCI-H522	> -4.00		
Colon Cancer	••••••		
COLO 205	> -4.00		
HCC-2998	> -4.00		
HCT-116	> -4.00		1
HCT-15	> -4.00		1
HT29	> -4.00		
KM12	> -4.00		
SW-620	> -4.00		l
CNS Cancer			
SF-268	> -4.00		į.
SF-295	> -4.00		1
SF-539	> -4.00		1
SNB-19	> -4.00		
U251	> -4.00		
Melanoma	- 4 M	*****************************	<u> </u>
· LOX IMVI	> -4.00 > -4.00		
M14 SK-MEL-2	> -4.00		
SK-MEL-28	> -4.00		
SK-MEL-28 SK-MEL-5	> 4.00		
UACC-257	> -4.00		
UACC-62	> -4.00		
Ovarian Cancer		***************************************	~
IGROVI	> -4.00		
OVCAR-3	> -4.00		
OVCAR-4	> -4.00		
OVCAR-5	> -4.00		i
OVCAR-8	> -4.00		
SK-OV-3	> -4.00		
Renal Cancer		***************************************	
786-0	> -4.00		-
A498	> -4.00		
ACHN	> -4.00		L
CAKI-1	-4.10 > -4.00		Γ .
RXF 393	> -4.00 > -4.00		
SNI2C TK-10	> -4.00		1
UO-31	> -4.00		[
Prostate Cancer	7.170		
PC-3	> -4.00		
DU-145	> -4.00		1
Breast Cancer	ļ	***************************************	
MCF7	> -4.00		
NCVADR-RES	> -4.00		1
MDA-MB-231/ATCC	> -4.00	•	
HS 578T	> -4.00		1
MDA-MB-435	> -4.00		
MDA-N	> -4.00		
BT-549	> -4.00		
T-47D	> -4.00		
			•
	1 4 444		1
MG_MID	-4.00		
Delta	0.10		į.
	l e		
Delta	0.10	<u> </u>	0 1 .2 .3

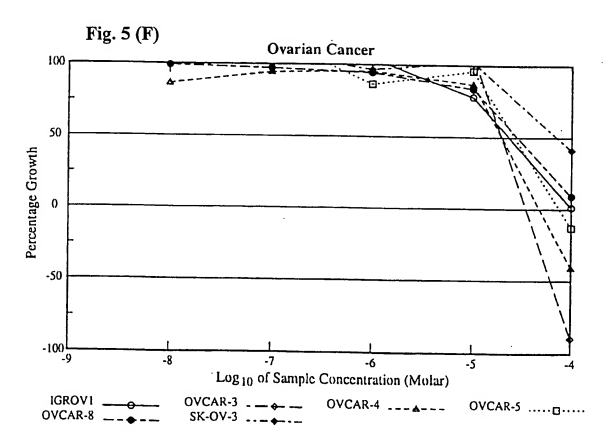


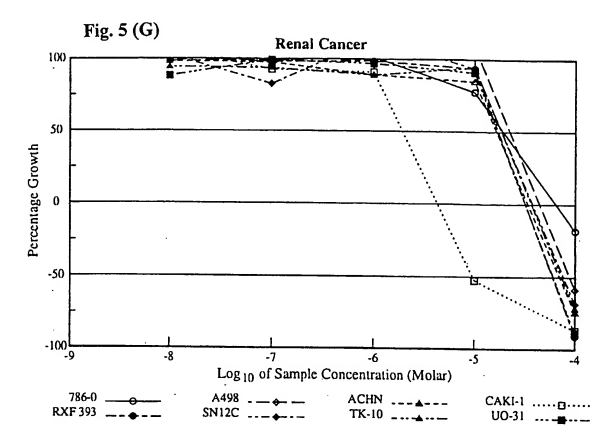


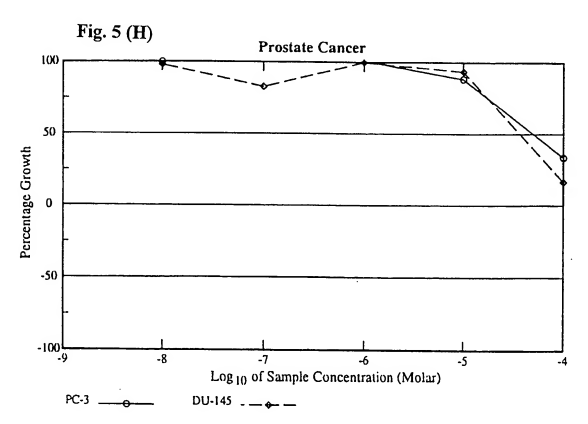












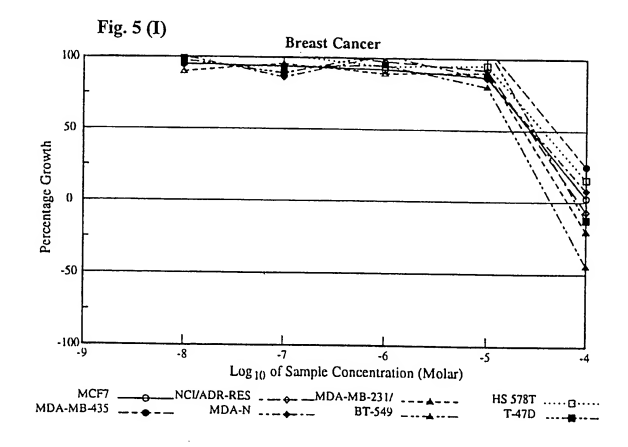
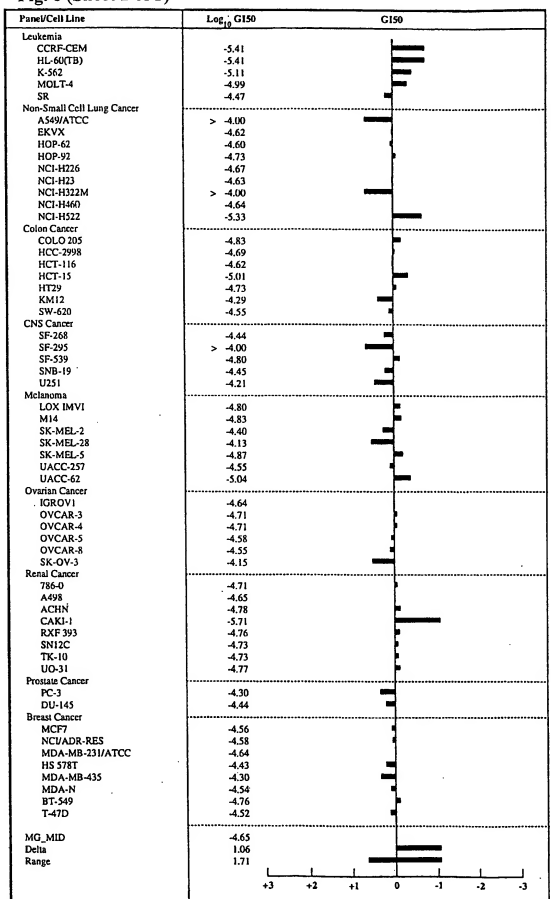


Fig. 6 (Sheet 1 of 3)

Panel/Cell Line	Log _{IO} TGI	TGI
Leukemia		
CCRF-CEM	-4.09	4
HL-60(TB)	-4.06	
K-562	-4.41	_
MOLT-4	> -4.00	
SR		3
	> -4.00	-
Non-Small Cell Lung Cancer		***************************************
AS49/ATCC	> -4.00	=
EKVX	-4.27)
HOP-62	-4.15	*
HOP-92	-4.38	j a
NCI-H226	-4.25	1
NCI-H23	-4.28	
NCI-H322M		
	> -4.00	7
NCI-H460	-4.19	i
NCI-H522	-4.61	pages.
Colon Cancer	******************************	
COLO 205	-4.49	—
HCC-2998	-4.34	•
HCT-116	4.13	Ţ
HCT-15		L
	-4.45	·
HT29	> -4.00	–
KM12	> -4.00	~
SW-620	> -4.00	=
CNS Cancer		•
SF-268	> -4.00	
SF-295	> -4.00	=
SF-539		٦
	-4.44	. F
SNB-19	> -4.00	4
U251	> -4.00	—
Melanoma		
LOX IMVI	-4.44	
M14 ·	-4.48	_
SK-MEL-2	> -4.00	
SK-MEL-28		3
	> -4.00	7
SK-MEL-5	-4.48	. F
UACC-257	-4.09	9
UACC-62	-4.59	
Ovarian Cancer	***************************************	
IGROVI	> -4.00	
OVCAR-3	-4.46	
OVCAR-4	-4.32	<u></u>
OVCAR-5	-4.12	<u>J</u>
OVCAR-8		_]
	> -4.00	
SK-OV-3	> -4.00	"
Renal Cancer	******************************	••••••
786-0	-4.19	(
A498	-4.35	 -
ACHN	-4.47	-
CAKI-I	-5.37	
RXF 393	-4.49	
SN12C		_
TK-10	-4.42	
	-4.43	.
UO-31	-4.49	. .
Prostate Cancer	***************************************	
PC-3	> -4.00	=
DU-145	> -4.00	=
Breast Cancer		
MCF7	> -4.00	
NCVADR-RES		7
	-4.07	7
MDA-MB-231/ATCC	-4.19	
HS 578T	> -4.00	=
MDA-MB-435	> -4.00	=
MDA-N	> -4.00	=
BT-549	-4.36	L
T-47D		_
1-410	-4.10	7
MC MID		***************************************
MG_MID	-4.22	
Delta	1.15	
Range	1.37	-
	·	<u> </u>
	+3 +	2 +1 0 -1 -2 -3

WO 98/49146

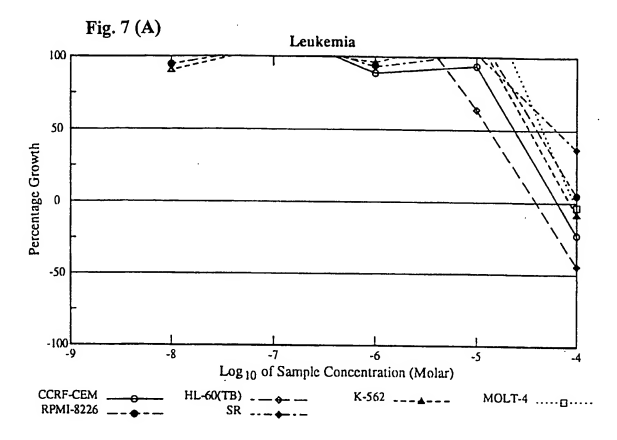
Fig. 6 (Sheet 2 of 3)

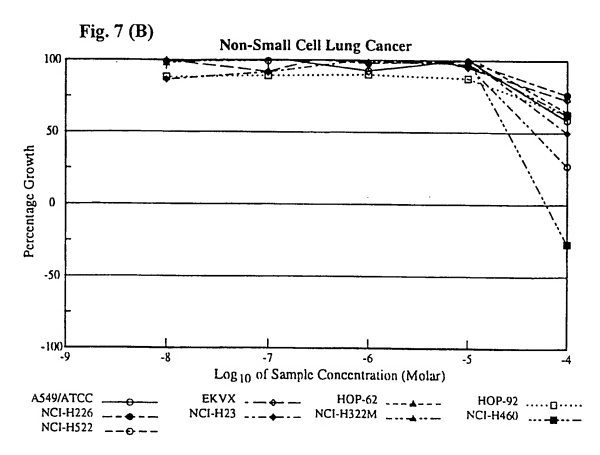


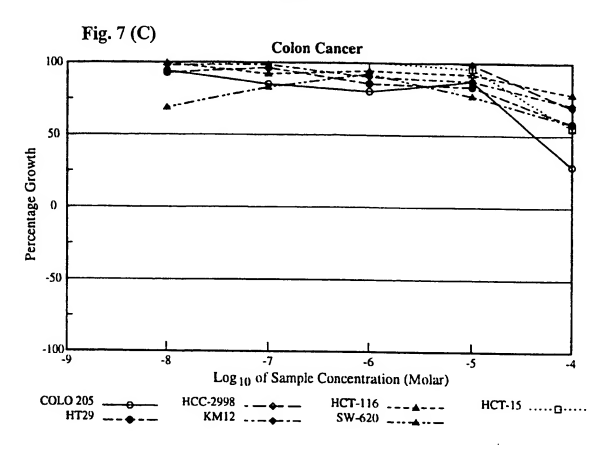
WO 98/49146 PCT/US98/08602

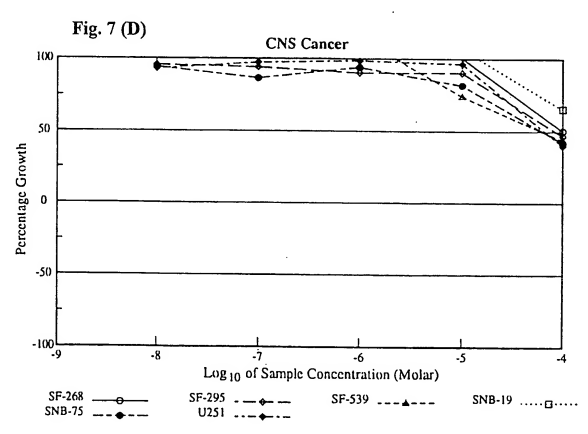
Fig. 6 (Sheet 3 of 3)

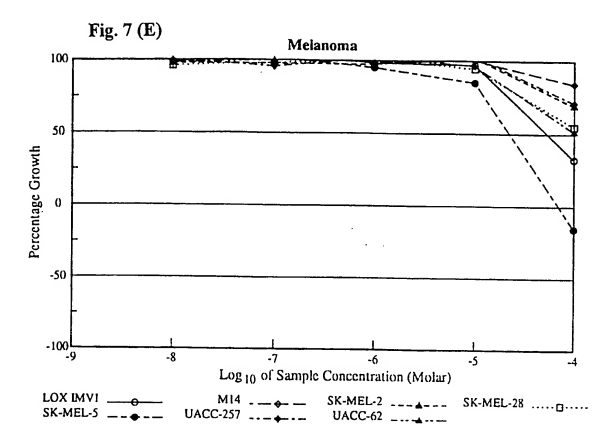
Leibenia CCRP CEM	Panel/Cell Line	Log ₁₀ LC50	LC50	
CCRF-CEM H. H. 607(TB) K. 562 MOLT-4 SR MOLT-4 SR Non-Small Cell Lung Cancer A549/ATCC EKVX A-4.00 HOP-92 A-4.00 HOP-92 NCI-H223 NCI-H223 NCI-H223 NCI-H224 NCI-H23 NCI-H322 NCI-H322 NCI-H322 NCI-H322 NCI-H322 NCI-H322 NCI-H322 NCI-H323 NCI-H322 A-4.00 NCI-H3460 NCI-H360 NCI-H360 NCI-H360 NCI-H360 NCI-H37 NCI-H360 NCI-H	Leukemia			
HL-60(TB)		> -4.00	1	
K. \$62		•		
MOLT-4 > -4.00				
SR			1	
Non-Small Cell Lung Cancer			46, 3, 40	
A549/ATCC EEVX HOP-62 HOP-62 HOP-92 A-00 NCI-H226 NCI-H23 NCI-H23 NCI-H322M NCI-H322M NCI-H322M NCI-H456 NCI-H322 Colon Cancer COLO 205 HCC-2908 HCT-116 HCT-15 HCT-19 HCT-15 HCT-19 KM12 SW-620 CNS Cancer SF-268 SF-295 SF-39 SR-95 SF-39 SR-95 SR-		> -4.00		
ERVX			•••••••••••••••••••••••••••••••••••••••	•••••••
HOP-62 HOP-92 HOP-92 NCI-H226 NCI-H23 NCI-H322M NCI-H322M NCI-H456 NCI-H322 NCI-H456 NCI-H322 NCI-H456 NCI-H522 -4.09 NCI-H522 -4.00 HCT-116 -4.00 HCT-15 -4.00 HCT-15 -4.00 SW-420 -4.00 SW-420 -4.00 SW-420 -4.00 SSF-268 SF-295 SF-539 -4.00 SF-268 SF-295 SF-539 -4.00 SF-539 SNB-19 -4.00 U251 -4.00 McI-M14 -4.14			4	
HOP-92	EKVX			
NCI-H226	HOP-62	> -4.00		
NCI-H232	HOP-92	-4.03		
NCI-H322M	NCI-H226	> -4.00	•	
NCI-H460 NCI-H522 A09 Colon Cancer COLO 205 HCC-2998 A-410 HCT-116 A-400 HCT-115 A-400 HTT29 A-400 KM12 SW-620 SW-620 SW-620 SF-268 SF-268 SF-295 SW-620 SF-295 SW-820 SF-399 A-400 SF-295 SW-819 A-400 McIanoma LDX IMVI A-4,14 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-3 SK-	NCI-H23	> -4.00	ţ .	
NCI-HS22 Colon Canner COLO 205 HCC-2998 HCT-116 > 4.00 HCT-15 > 4.00 HCT-15 > 4.00 SW-620 SW-620 SW-620 CNS Cancer SF-206 SF-295 SF-399 SNB-19 LOX IMVI M41 SK-MEL-2 SK-MEL-2 SK-MEL-5 UACC-257 UACC-62 SK-MEL-5 UACC-257 UACC-62 SK-MEL-5 UACC-83 SK-MEL-5 UACC-84 SK-MEL-8 SK-MEL-5 UACC-257 SUACC-82 SK-MEL-5 SK-M	NCI-H322M	> -4.00	•	
NCI-HS22 Colon Canner COLO 205 HCC-2998 HCT-116 > 4.00 HCT-15 > 4.00 HCT-15 > 4.00 SW-620 SW-620 SW-620 CNS Cancer SF-206 SF-295 SF-399 SNB-19 LOX IMVI M41 SK-MEL-2 SK-MEL-2 SK-MEL-5 UACC-257 UACC-62 SK-MEL-5 UACC-257 UACC-62 SK-MEL-5 UACC-83 SK-MEL-5 UACC-84 SK-MEL-8 SK-MEL-5 UACC-257 SUACC-82 SK-MEL-5 SK-M	NCI-H460	> -4.00		
Colon Cancer				
COLO 205 HC2-998 HC7-116 COLO 205 HC7-1298 HC7-116 COLO 205 HC7-116 HC7-15 S - 4.00 HC7-116 HC7-15 S - 4.00 KM12 SW-620 SW-620 SW-620 CNS Cancer SF-268 SF-295 SF-399 SNB-19 U251 Mclanoma LOX IMVI 4.07 Ml4 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-3 SK-MEL-3 SK-MEL-3 UACC-257 UACC-62 UACC-62 Vuarian Cancer IOROVI OVCAR-3 OVCAR-3 OVCAR-3 SK-OV-3 S		***************************************	***************************************	
HCC-2998 HCT-16 HCT-15 HCT-16 HCT-16 HCT-16 HCT-15 HCT-16		-4.15		
HCT-116 HCT-15 HCT-15 HCT-15 HCT-15 HCT-15 HCT-19 XM12 XW-620 XW-620 XW-620 XW-620 CNS Cancer SF-268 SF-295 SF-399 SR-399 SR-19 U251 Mclanoma LOX IMVI M14 XK-MEL-2 XK-MEL-2 XK-MEL-2 XK-MEL-2 XK-MEL-2 XK-MEL-2 XK-MEL-2 XK-MEL-3 X			[
HCT-15			1	
HT29			1	
KM12 SW-620 CNS Cancer SF-268 SF-295 SF-295 SF-399 SNB-19 U2S1 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-28 SK-MEL-28 SK-MEL-5 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 SW-OVCAR-5 SK-OV-3 Renal Cancer 786-0 A498 ACHN A15 CAKI-1 CAKI-1 SK-MSH CAKI-1 SK-MSH SW-OVCAR-1 SW			}	
SW-620 CNS Cancer SF-268 SF-268 SF-295 SF-539 -4.00 SNB-19 U251 Melanoma LOX IMVI M14 -4.14 SK-MEL-2 SK-MEL-2 SK-MEL-28 SK-MEL-5 UACC-257 UACC-257 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-8 SK-OV-3 SK-O			j	
CNS Cancer SF-268 SF-295 SF-295 SN-19 U251 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-28 SK-MEL-28 SK-MEL-38 SK-MEL-36 SK-MEL-37 UACC-62 UACC-62 UACC-62 UACC-62 UACC-62 UACC-63 UACC-63 OVCAR-3 OVCAR-3 OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 SK-11 SK-11 SK-11 SK-11 SK-12 SK-11 SK	•		3	
SF-268 SF-295 SF-399 SNB-19 U251 V251 V251 V251 V3-400 Melanoma LOX IMVI M14 SK-MEL-2 SK-MEL-2 SK-MEL-2 SK-MEL-5 V3-400 V3-400 V3-400 V4-400 V		> -4.00	<u> </u>	
SF-295 SF-539 SNB-19 U251 Melanoma LOX IMV1 M14 SK-MEL-28 SK-MEL-28 SK-MEL-5 UACC-257 UACC-62 Ovarian Cancer IGROV1 OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 COX-1 COX			***************************************	******
SF-339 SNB-19 U251 Melanoma LOX IMV1 M14 SK-MEL-2 SK-MEL-22 SK-MEL-28 SK-MEL-5 UACC-257 UACC-62 UACC-267 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-4 OVCAR-8 SK-W-3 SK-W-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAMI CAKI-1 CAMI CAKI-1 CAMI CAKI-1 CAMI CAKI-1 CAMI CAMI CAKI-1 CAMI CAMI CAMI CAMI CAMI CAMI CAMI CAMI			•	
SNB-19			•	
U251			}	
Mclanoma	· · · · · · · · · · · · · · · · · · ·		\$	
LOX IMVI MI4 SK-MEL-2 SK-MEL-28 SK-MEL-5 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-4 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 CAKI-1 CAKI-1 COKI-1 TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-23I/ATCC HS 5787 MDA-MB-435 MDA-MB-435 MDA-MB BT-549 T-47D MG_MID Delta MG_MID Delta A400 A400 A400 A400 A400 A400 A400	U251	> -4.00	•	
MI4 SK-MEL-2 SK-MEL-2 SK-MEL-5 UACC-257 UACC-257 VACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 CAKI-1 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCIVADR-RES MDA-MB-231/ATCC HS 578T Delta MG_MID MG_MID MG_MID MG_MID Delta -4.00 -4.0	Melanoma			•••••
SK-MEL-2 SK-MEL-28 SK-MEL-5 UACC-57 UACC-62 Ovrian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 CAKI-1 TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-MB BT-549 T-47D MG_MID MG_MID MG_MID MG_MID MG_MID Delta	LOX IMVI	-4.07		
SK-MEL-28 SK-MEL-3 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 SN12C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breas Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-M BT-549 T-47D MG_MID MG_MID MG_MID MG_MID MG_MID Delta - 4.00 -4.00 -4.00 -4	M14	-4.14)	
SK-MEL-28 SK-MEL-3 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 SN12C TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breas Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-M BT-549 T-47D MG_MID MG_MID MG_MID MG_MID MG_MID Delta - 4.00 -4.00 -4.00 -4	SK-MEL-2	> -4.00	(
SK-MEL-5 UACC-257 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 CAKI-1 TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breas Cancer MCF7 NCUADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID MG_MID Delta -4.05 Delta -4.00 -4.16 -4.00 -4.16 -4.00 -4.05 -4.00 -4.00 -4.00 -4.00 -4.00 -4.05 -4.00			i	
UACC-257 UACC-62 UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 CAKI-1 TK-10 UO-31 Prostate Cancer PC-2 PC-2 PC-2 PC-3 DU-145 Breast Cancer MCF7 NCU/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta > 4.00			. i	
UACC-62 Ovarian Cancer IGROVI OVCAR-3 OVCAR-4 OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 CAKI-1 TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCUADR-RES MDDA-MB-23I/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta > 4.00			· •	
Ovarian Cancer IGROVI > -4.00 OVCAR-3 -4.20 -4.00 OVCAR-4 > -4.00 -4.00 OVCAR-5 > -4.00 -4.00 SK-OV-3 > -4.00 -4.00 Renal Cancer 786-0 > -4.00 A498 -4.05 -4.05 ACHN -4.15 -5.02 RXF 393 -4.22 -4.11 SN12C -4.11 -4.13 UO-31 -4.21 -4.21 Prostate Cancer PC-3 > -4.00 DU-145 > -4.00 -4.00 Breast Cancer -4.00 -4.00 MCF7 > -4.00 -4.00 MDA-MB-231/ATCC > -4.00 -4.00 HS 578T > -4.00 -4.00 MDA-MB-35 > -4.00 -4.00 MDA-N > -4.00 -4.00 BT-549 > -4.00 -4.00 T-47D > -4.00 -4.05 MG_MID -4.05 -4.05 Delta 0.97			L	
IGROVI	•	***************************************		
OVCAR-3 -4.20 OVCAR-4 > -4.00 OVCAR-5 > -4.00 OVCAR-8 > -4.00 SK-OV-3 > -4.00 Renal Cancer -4.05 786-0 > -4.00 A498 -4.05 ACHN -4.15 CAKI-1 -5.02 RXF 393 -4.22 SN12C -4.11 TK-10 -4.13 UO-31 -4.21 Prostate Cancer PC-3 PC-3 > -4.00 DU-145 > -4.00 Breast Cancer -4.00 MCF7 > -4.00 NCI/ADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-N > -4.00 MDA-N > -4.00 MDA-N > -4.00 MT-47D > -4.00 MG_MID -4.05 Delta 0.97		> -4 (V)		
OVCAR-4 > -4.00 OVCAR-5 > -4.00 OVCAR-8 > -4.00 SK-OV-3 > -4.00 Renal Cancer -4.05 786-0 -4.05 ACHN -4.15 CAKI-1 -5.02 RXF 393 -4.22 SN12C -4.11 TK-10 -4.13 UO-31 -4.21 Prostate Cancer -4.00 PC-3 > -4.00 DU-145 > -4.00 Breast Cancer -4.00 MCF7 > -4.00 NCI/ADR-RES -4.00 MDA-MB-23I/ATCC -4.00 HS 578T > -4.00 MDA-N -4.00 BT-549 -4.00 T-47D -4.00 MG_MID -4.05 Delta 0.97			L	
OVCAR-5 OVCAR-8 SK-OV-3 Renal Cancer 786-0 A498 ACHN CAKI-1 CAKI-1 TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta > 4.00 > 4.00 > 4.00 > 4.00 > 4.00 > 4.00 > 4.00 4.00 6.00 6.00 6.00 6.00 7.400 7.470 7.400			₽	
OVCAR-8 SK-OV-3 > 4.00 Renal Cancer > 4.00 786-0 A498 -4.05 ACHN -4.15 CAKI-1 CAKI-1 -5.02 RXF 393 -4.22 SN12C -4.11 TK-10 -4.13 UO-31 -4.21 Prostate Cancer > -4.00 DU-145 > -4.00 Breast Cancer > -4.00 MCF7 > -4.00 NCI/ADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-MB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97			}	
SK-OV-3			}	
Renal Cancer 786-0 A498 ACHN	· · · · · · · · · · · · · · · · · · ·		3	
786-0 > -4.00 A498 -4.05 ACHN -4.15 CAKI-1 -5.02 RXF 393 -4.22 SN12C -4.11 TK-10 -4.13 U0-31 -4.21 Prostate Cancer > -4.00 DU-145 > -4.00 Breast Cancer > -4.00 MCF7 > -4.00 NCVADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-NB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97		> -4.00	1	
A498 ACHN CAKI-1		4 1/1	······	
ACHN CAKI-1 CAKI			j	
CAKI-1 RXF 393 -4.22 SN12C TK-10 UO-31 -4.13 UO-31 -4.21 Prostate Cancer PC-3 DU-145 -4.00 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta -5.02 -4.22 -4.11 -4.22 -4.11 -4.13 -4.21 -4.00 -4.01 -4.00 -4.05 -4.00 -4.00 -4.05 -4.00			L	
RXF 393 SN12C TK-10 UO-31 VO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta -4.22 -4.11 -4.12 -4.11 -4.13 -4.21 -4.00 -4.00			<u> </u>	
SN12C TK-10 TK-10 UO-31 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta -4.11 -4.13 -4.21 -4.00 > -4.00 > -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00				
TK-10 UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta -4.13 -4.21 -4.00 > -4.00 > -4.00 > -4.00 > -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00 -4.00			.	
UO-31 Prostate Cancer PC-3 DU-145 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta -4.00 -4.01 -4.01 -4.01 -4.01 -4.01 -4.05 -4.00 -4.05 -4.00 -4.05 -4.00 -4.05 -4.05 -4.00 -4.05 -4.05 -4.05 -4.05 -4.07 -4.05 -4.07 -4.08 -4.09 -4.09 -4.09 -4.09			7	
Prostate Cancer > -4.00 DU-145 > -4.00 Breast Cancer > -4.00 MCF7 > -4.00 NCI/ADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-MB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97			ř	
PC-3 DU-145 Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta > -4.00 > -4.00 > -4.00 MG_MID Delta > -4.05 > -4.00 > -4.05 Delta		-4.21	P	
DU-145 > -4.00 Breast Cancer -4.00 MCF7 > -4.00 NCI/ADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-MB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97			***************************************	
Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta MG_MID Delta S-4.00 -4.05 -4.05 -6.00 -4.05 -6.07 -4.05 -6.00 -4.05 -6.07 -4.05 -6.00 -4.05 -6.07	PC-3	> -4.00	•	
Breast Cancer MCF7 NCI/ADR-RES MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta MG_MID Delta S-4.00 -4.05 -4.05 -6.00 -4.05 -6.07 -4.05 -6.00 -4.05 -6.07 -4.05 -6.00 -4.05 -6.07	DU-145	> -4.00	•	
MCF7 > -4.00 NCVADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-MB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97				·
NCI/ADR-RES > -4.00 MDA-MB-231/ATCC > -4.00 HS 578T > -4.00 MDA-MB-435 > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97		> -4.00	1	
MDA-MB-231/ATCC				
HS 578T > -4.00 MDA-MB-435 > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97			ì	
MDA-MB-435 > -4.00 MDA-N > -4.00 BT-549 > -4.00 T-47D > -4.00 MG_MID -4.05 Delta 0.97			l l	
MDA-N BT-549 T-47D MG_MID Delta MDA-N S - 4.00 S - 4.00 S - 4.00 S - 4.00 S - 4.05			1	
BT-549 > -4.00 > -4.00 MG_MID -4.05 Delta 0.97			. 1	
T-47D > -4.00 MG_MID			1	
MG_MID -4.05 Delta 0.97			1	
Delta 0.97	1-4/0	> -4.00	1	
Delta 0.97	14G 14TD :	**************************************		••••••
Kange 1.02				
	Kange .	1.02		
				
+3 +2 +1 0 -1 -2		+3	+2 +1 0 -1 -2	-3
		i		

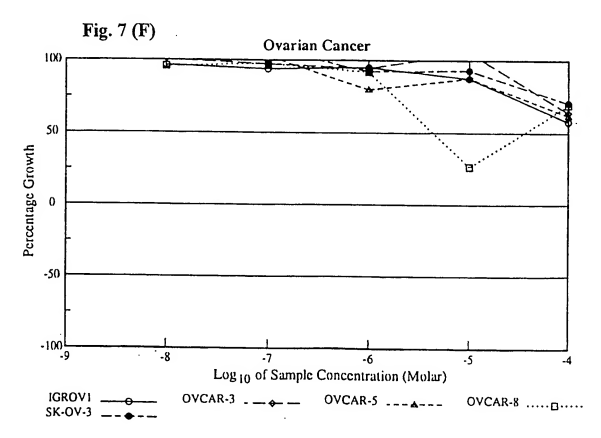


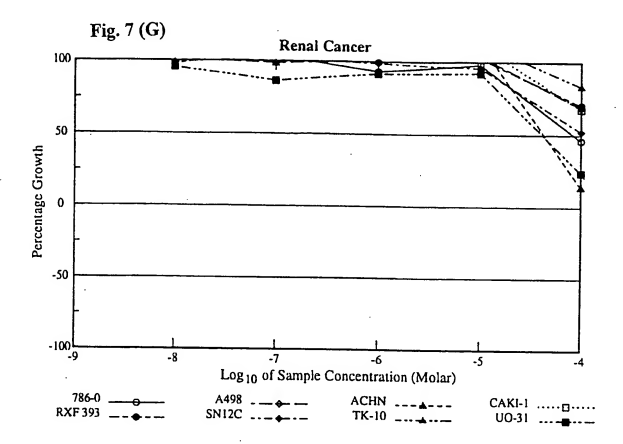


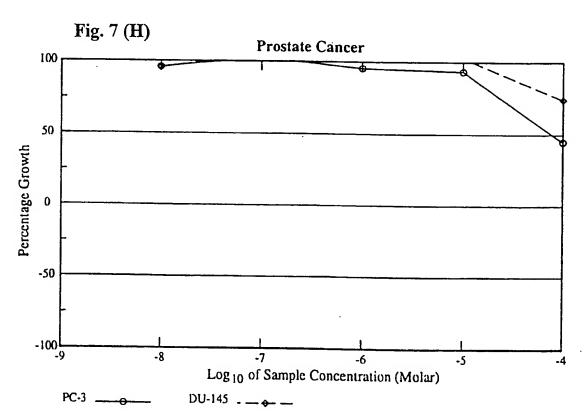


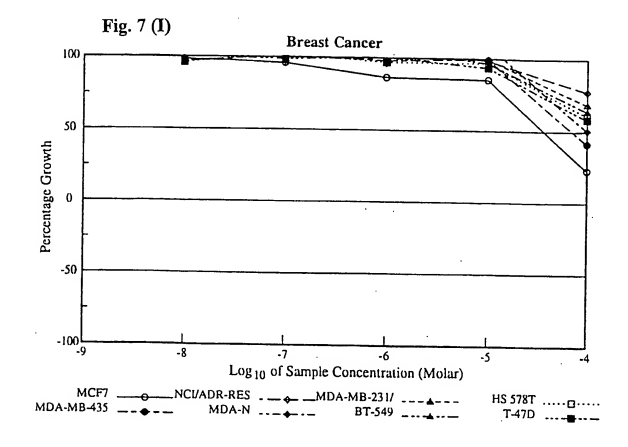












WO 98/49146 PCT/US98/08602

Fig. 8 (Sheet 1 of 3)

Panel/Cell Line	Log ₁₀ G150	G150
Leukemia		
CCRF-CEM	-4.63	
HL-60(TB)	-4.87	
K-562	-4.48	
MOLT-4	-4.34	
RPMI-8226	-4.40	<u> </u>
SR		Γ
	-4.20	
Non-Small Cell Lung Cancer		***************************************
A549/ATCC	> -4.00	–
EKVX	> -4.00	4
HOP-62	> -4.00	4
HOP-92	> -4.00	=
NCI-H226	> -4.00	■
NCI-H23	-4:00	₫
NCI-H322M	> -4.00	<u>.</u>
NCI-H460	-4.60	7
NCI-H522		
	-4.33	_
Colon Cancer		***************************************
COLO 205	4.26	P
HCC-2998	> -4.00	=
HCT-116	-4.20	•
HCT-15	-4.61	—
HT29	-4.25	L Company
KM12	> 4.00	
SW-620	4.24	l,
CNS Cancer	7,24	<u></u>
SF-268	_ A00	
	> -4.00	3
SF-295	4.07	
SF-539	-4.20)
SNB-19	> -4.00	•
SNB-75	-4.20)
U251	-4.18	}
Melanoma		***************************************
LOX IMVI	-4.27	
M14	> -4.00	
SK-MEL-2	> -4.00	3
SK-MEL-28	> 4.00	3
SK-MEL-5	•	
UACC-257	-4.65	
	> -4.00	4
UACC-62	> -4.00	7
Ovarian Cancer	***************************************	
IGROVI	> -4.00	=
OVCAR-3	> -4.00	
OVCAR-5	> -4.00	
OVCAR-8		Į.
SK-OV-3	> -4.00	<u>.</u>
Renal Cancer	***************************************	
786-0	-4.07	***************************************
A498	> 4.00	j
ACHN		1
CAKI-I	-4.37	_
	> 4.00	5
RXF 393	> -4.00	•
SN12C	> -4.00	•
TK-10	> -4.00	=
UO-31	-4.38	>
Prostate Cancer		
PC-3	-4.11	
DU-145	> -4.00	.
Breast Cancer		1
MCF7	-4.43	
	,	<u></u>
NCI/ADR-REC	> 4.00 > 4.00	3
NCVADR-RES		–
MDA-MB-231/ATCC		
MDA-MB-231/ATCC HS 578T	> -4.00	=
MDA-MB-231/ATCC HS 578T MDA-MB-435	> -4.00 -4.15	•
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N	> -4.00	1
MDA-MB-231/ATCC HS 578T MDA-MB-435	> -4.00 -4.15	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N	> -4.00 -4.15 > -4.00 > -4.00	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549	> 4.00 -4.15 > -4.00	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D	> -4.00 -4.15 > -4.00 > -4.00 > -4.00	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID	> -4.00 -4.15 > -4.00 > -4.00 > -4.00	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta	> 4.00 -4.15 > -4.00 > -4.00 > -4.00	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID	> -4.00 -4.15 > -4.00 > -4.00 > -4.00	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta	> -4.00 -4.15 > -4.00 > -4.00 > -4.00 -4.15 0.72 0.87	
MDA-MB-231/ATCC HS 578T MDA-MB-435 MDA-N BT-549 T-47D MG_MID Delta	> 4.00 -4.15 > -4.00 > -4.00 > -4.00	2 +1 01 -2 -3

Fig. 8 (Sheet 2 of 3)

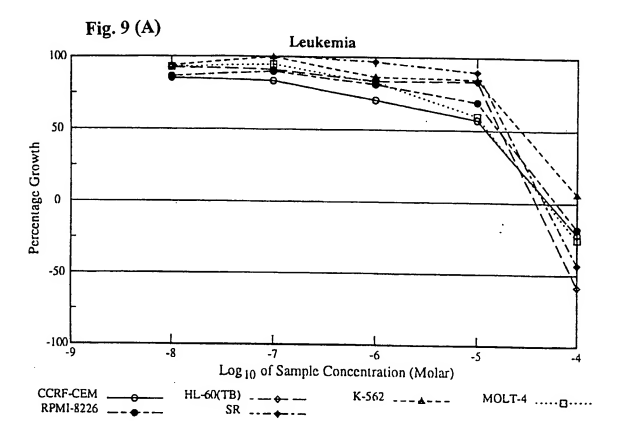
Panel/Cell Line	Log ₁₀ TGI	TGI	
Leukemia	1	· · · · · · · · · · · · · · · · · · ·	
CCRF-CEM	-4.19)	
HL-60(TB)	-4.41		
K-562	-4.07	•	
MOLT-4	-4.02	i	
RPMI-8226	> -4.00	į	
SR	> -4.00	į	
Non-Small Cell Lung Cancer			*****
AS49/ATCC	> -4.00		
EKVX	> -4.00	l l	
HOP-62	> -4.00	.	
HOP-92	> -4.00		
NCI-H226	> 4.00		
NCI-H23	> -4.00		
NCI-H322M	> -4.00		
NCI-H460	-4.21	 	
NCI-H522	> -4.00	•	
Colon Cancer			******
COLO 205	> -4.00		
HCC-2998	> -4.00	į.	
HCT-116	> -4.00	ł	
HCT-15	-4.07	,	
HT29	> -4.00	ĺ	
KM12	> -4.00	1	
SW-620	> -4.00	1	
CNS Cancer			
SF-268	> -4.00		
SF-295	> -4.00		
SF-539	> -4.00	1	
SNB-19	> -4.00	1	
SNB-75	> -4.00		
U251	> -4.00	1	
Melanoma	,		
LOX IMVI	> -4.00		***************************************
M14	> -4.00	l	
SK-MEL-2	> -4.00		
SK-MEL-28	> -4.00	1	
SK-MEL-5	-4.15	(L	
UACC-257	> -4.00	Γ	
UACC-62	> -4.00	<i>y</i>	
Ovarian Cancer	***************************************		
IGROV1	> -4.00		***********************
OVCAR-3	> 4.00		
OVCAR-5	> -4.00	1	
OVCAR-8	> -4.00		
SK-OV-3	> 4.00	1	
Renal Cancer	7-7.00		,
786-0	> -4.00		***************************************
A498	> 4.00	All I	
ACHN	> 4.00		
CAKI-I	> 4.00	A I	
RXF 393	> 4.00	1	
SNI2C	> 4.00		
TK-10	> 4.00	1	
UO-31	> 4.00		
Prostate Cancer	7.00	THE	
PC-3	> -4.00	1	•••••••
DU-145	> -4.00	I	
Breast Cancer	7		
MCF7	> -4.00		
NCI/ADR-RES	> -4.00		
MDA-MB-231/ATCC			
HS 578T	> -4.00		1
MDA-MB-435	> -4.00		
MDA-N	> -4.00	Y	
BT-549	> -4.00	A .	
	> -4.00		
T_47D	> -4.00		i
T-47D			
MG_MID	-4.02	***************************************	
MG_MID Delia	0.39		
MG_MID			•••••••••••••••••••••••••••••••••••••••
MG_MID Delta	0.39 0.41		
MG_MID Delta	0.39 0.41	2 +1 0 -1	-2 -3

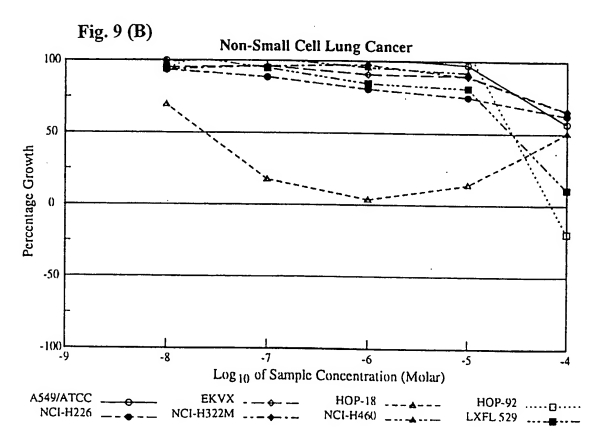
Fig. 8 (Sheet 3 of 3)

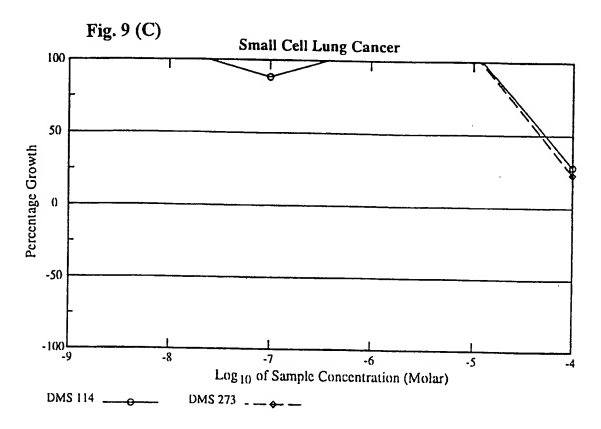
32/47

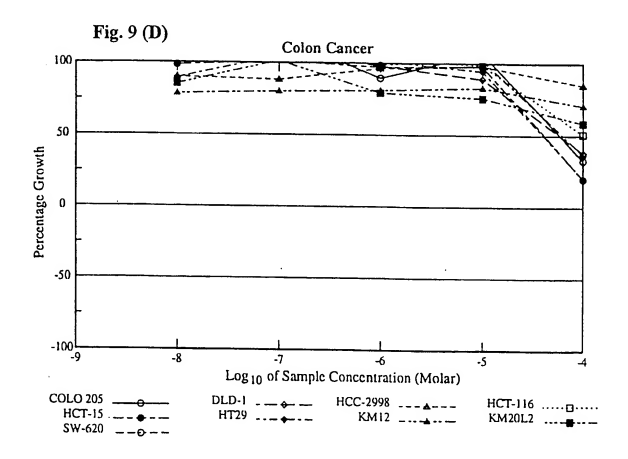
Panel/Cell Line	Log ₁₀ LC50	LC50
Leukemia	1	T .
CCRF-CEM	> -4.00	1
HL-60(TB)	> -4.00	1
K-562	> -4.00	i
MOLT-4	. > -4.00	i
RPMI-8226	> -4.00	i i
SR	> -4.00	
Non-Small Cell Lung Cancer	***************************************	
A549/ATCC	> -4.00	
EKVX	> -4.00	
HOP-62	> -4.00	· ·
HOP-92	> -4.00	
NCI-H226	> -4.00	
NCI-H23	> -4.00	
NCI-H322M	> -4.00	
NCI-H460	> -4.00	
NCI-H522	> -4.00	
Colon Cancer		
COLO 205	> -4.00	
HCC-2998	> -4.00	
HCT-116	> -4.00	İ
HCT-15	> -4.00	
HT29	> -4.00	
KM12	> -4.00	1
SW-620	> -4.00	} ·
CNS Cancer	***************************************	
SF-268	> -4.00	}
SF-295	> -4.00	
SF-539	> -4.00	
SNB-19	> -4.00	•
SNB-75	> -4.00	į
U251	> -4.00	1
Melanoma	***************************************	***************************************
LOX IMVI	> -4.00	
M14 ·	> -4.00	
SK-MEL-2	> -4.00	
SK-MEL-28	> -4.00	
SK-MEL-5	> -4.00	
UACC-257	> -4.00	1
UACC-62	> -4.00	<u>}</u>
Ovarian Cancer	7.00	
IGROVI	> -4.00	
OVCAR-3	> -4.00	
OVCAR-5	> 4.00	
OVCAR-8	> 4.00	į
SK-OV-3	> 4.00	
	> 4.00	
Renal Cancer 786-0	> -4.00	. [
A498	> 4.00	}
ACHN	> 4.00]
CAKI-I	> 4.00	1
	> 4.00	1
RXF 393	> 4.00	1
SN12C	> 4.00	
TK-10		1
UO-31	> -4.00	
Prostate Cancer	> 400	
PC-3	> 4.00	
DU-145	> -4.00	
Breast Cancer	. 400	
MCF7	> 4.00	
NCVADR-RES	> -4.00	
MDA-MB-231/ATCC	> -4.00	
HS 578T	> -4.00	
MDA-MB-435	> -4.00	1
MDA-N	> -4.00	
BT-549	> -4.00	
T-47D	> -4.00	

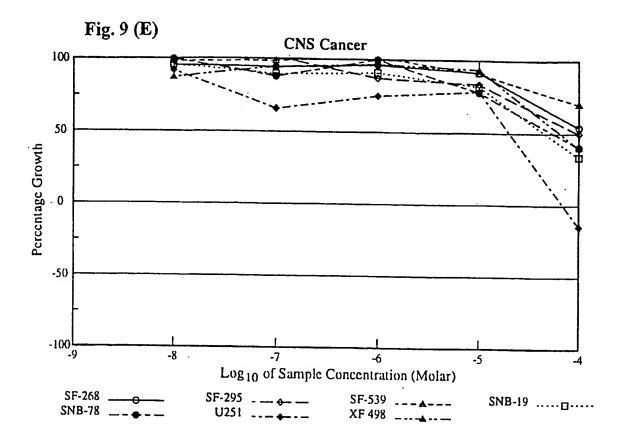
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Range	0.00	
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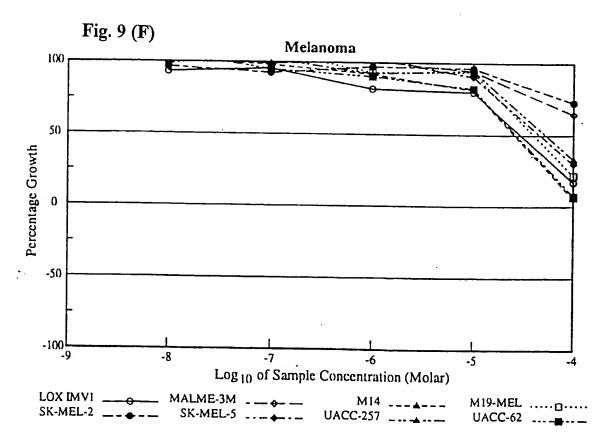


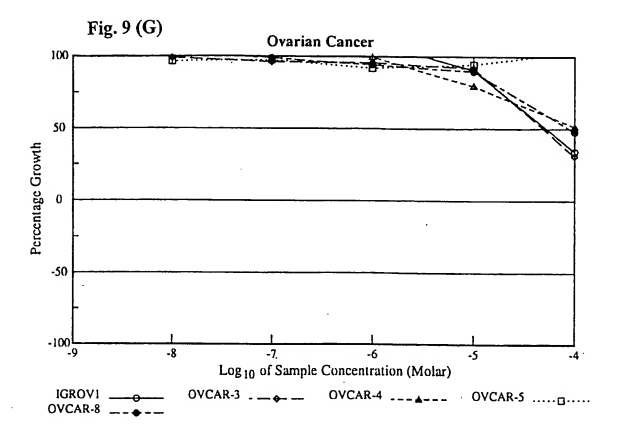












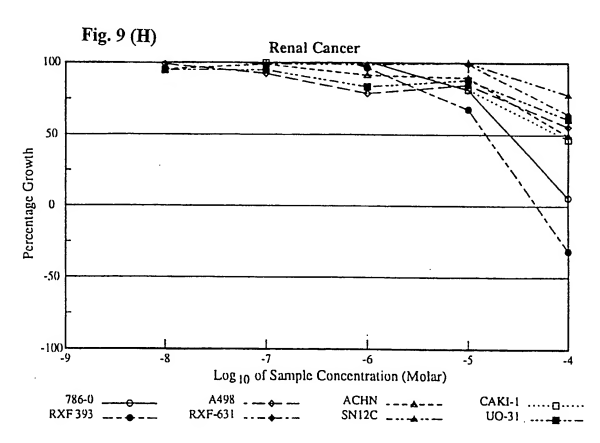


Fig. 10 (Sheet 1 of 3)

Panel/Cell Line	Log ₁₀ GI50	G150
Leukemia		1
CCRF-CEM	-4.91	
HL-60(TB)	-4.76	
K-562	-4.56	
MOLT-4	-4.89	
RPMI-8226	-4.78	
SR	-4.70	
Non-Small Cell Lung Cancer	***************************************	
A549/ATCC	> -4.00	
EKVX	> -4.00	
HOP-18		
HOP-92	-4.55	
NCI-H226	> -4.00	
NCI-H322M	> -4.00	
NCI-H460		
LXFL 529	-4.56	
Small Cell Lung Cancer	4.50	
DMS 114	-4.28	
DMS 273		į
	-4.33	V . □
Colon Cancer	4.00	······································
COLO 205	-4.23]
DLD-1	-4.23	
HCC-2998	> -4.00	
HCT-116	> -4.00	
HCT-15	-4.39	•
HT29	-4.40)•
KM12	> -4.00	=
KM20L2	> -4.00	=
SW-620	-4.20	•
CNS Cancer		
SF-268	> -4.00	
SF-295	· -4.00	
SF-539	> -4.00	
SNB-19	-4.34)
SNB-78	-4.25	
U251	-4.69	
XF 498	4.19	
Melanoma	***************************************	
LOX IMVI	-4.53	
MALME-3M	> -4.00	_
M14	-4.59	
M19-MEL	-4.39	
SK-MEL-2	> -4.00	
SK-MEL-5	-4.33	٦,
UACC-257	-4.33 -4.27	f
UACC-62	•	L
Ovarian Cancer	-4.57	
	4.00	***************************************
IGROVI	-4.28	Į.
OVCAR-3	-4.31	
OVCAR-4	> -4.00	
OVCAR-5	> -4.00	5
OVCAR-8	-4.05	-
Renal Cancer		
786-0	-4.59	
A498	> -4.00	
ACHN	-4.02	=
CAKI-I	-4.10	=
RXF 393	-4.82	—
RXF-631	> -4.00	-
SN12C	> -4.00	=
UO-31	> -4.00	
55 5 .	7 -7.00	
MG_MID	-4.28	
Delta		
Range	0.63	
raige	0.91	 , , , ,
	+3 +2	+1 0 -1 -2 -3

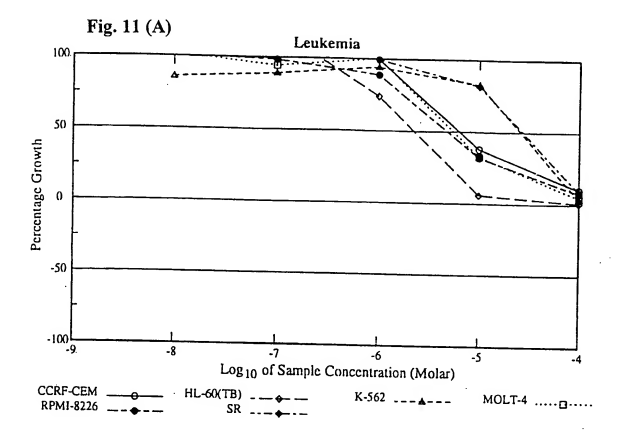
Fig. 10 (Sheet 2 of 3)

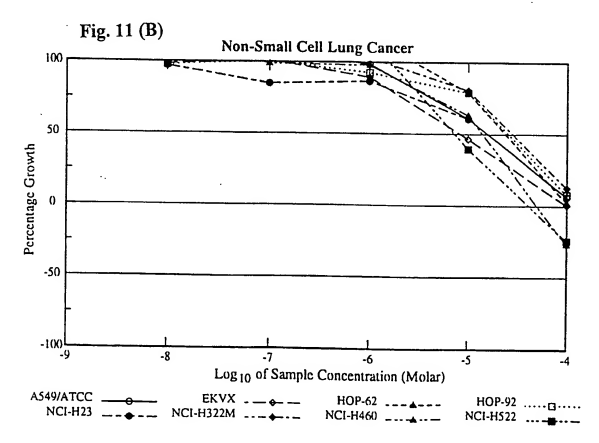
Panel/Cell Line	Log ₁₀ TGI	TGI	
Leukemia		Table Committee	
CCRF-CEM	· -4.28		
HL-60(TB)	-4.41 .		
K-562	> -4.00		
MOLT-4	-4.30		
RPMI-8226	-4.21		
SR	4.33		
Non-Small Cell Lung Cancer		••••••	
A549/ATCC	> -4.00	•	
EKVX	> -4.00	•	
HOP-18	> -4.00	¥.	
HOP-92	-4.16		
NCI-H226	> -4.00		
NCI-H322M	> -4.00	(1)	
NCI-H460			
LXFL 529	> -4.00	1	
Small Cell Lung Cancer	***************************************	***************************************	
DMS 114	> -4.00		
DMS 273	> -4.00		
Colon Cancer			***********
COLO 205	> -4.00		
DLD-I	> -4.00	Ì	
HCC-2998	> 4.00	}	
HCT-116	> -4.00		
HCT-15	> -4.00	}	
HT29	> 4.00	}	
KM12	> -4.00	Į.	
KM20L2	> -4.00	1	
SW-620	> -4.00	1	
CNS Cancer	7		
SF-268	> -4.00		
SF-295	> -4.00]	
SF-539	> 4.00]	
SNB-19	> -4.00	}	
	> -4.00	1	
SNB-78 U251	-4.16	L	
XF 498	> -4.00	ŗ	
Melanoma	> -4.00		
LOX IMVI	> -4.00		
	> 4.00]	
MALME-3M	1	1	
M14	> -4.00		
M19-MEL	> -4.00	1	
SK-MEL-2	> -4.00	}	
SK-MEL-5	> -4.00	1	
UACC-257	> -4.00	j	
UACC-62	> -4.00	•	
Ovarian Cancer	4.00	•••••	
IGROVI	> -4.00	5	
OVCAR-3	> -4.00	5	
OVCAR-4	> -4.00	1	
OVCAR-5	> -4.00	<u> </u>	
OVCAR-8	> -4.00	1	
Renal Cancer		•••••••••••••••••••••••••••••••••••••••	• • • • • • • • • • • • • • • • • • • •
786-0	> -4.00		
A498	> -4.00	- 24P	
ACHN	> -4.00		
CAKI-I	> -4.00		
RXF 393	-4.32		
RXF-631	> -4.00		
SN12C	> -4.00		
UO-31	> -4.00	4	
MG_MID	-4.04		
Delta	0.37	—	
Range	0.41	· -	
-	<u> </u>	1 1	
	+3 +2	+1 0 -1 -	2 -3
		•	_

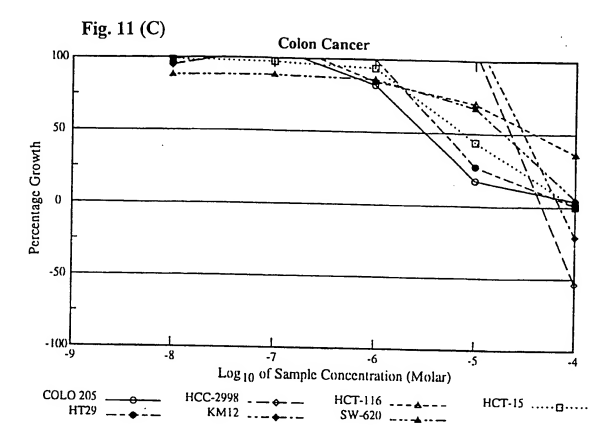
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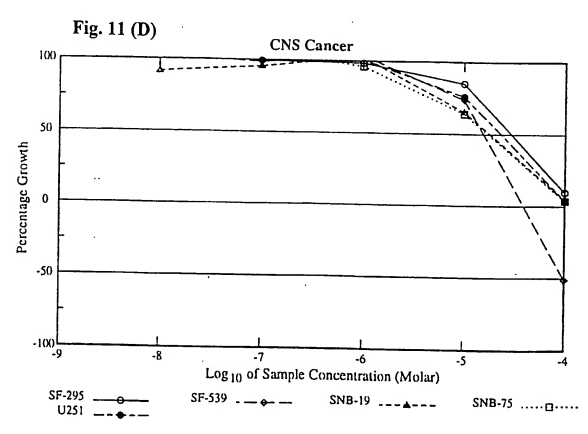
Fig. 10 (Sheet 3 of 3)

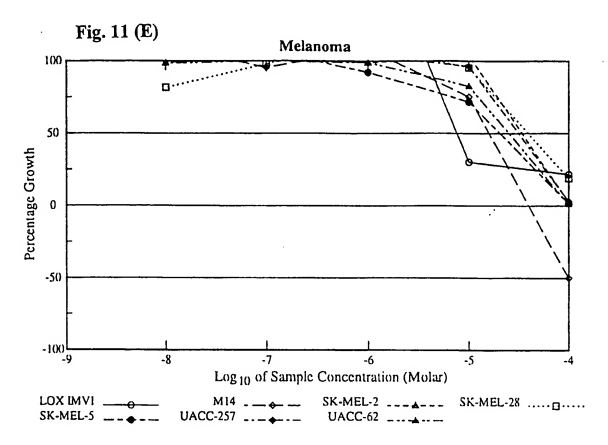
Panel/Cell Line	Log ₁₀ LC50	LC50
Leukemia		1
CCRF-CEM	> -4.00	1
HL-60(TB)	-4.06	1 1
K-562	> -4.00	1
MOLT-4	> -4.00	1 6
RPMI-8226	> -4.00	(P)
SR	> -4.00	
Non-Small Cell Lung Cancer		
A549/ATCC	> -4.00	
EKVX	> -4.00	
HOP-18	> -4.00	
HOP-92	> -4.00	
NCI-H226	> -4.00	
NCI-H322M	> -4.00	
NCI-H460		
LXFL 529	> -4.00	
Small Cell Lung Cancer		
DMS 114	> -4.00	
DMS 273	> -4.00	
Colon Cancer		
COLO 205	> -4.00	
DLD-1	> -4.00	
HCC-2998	> -4.00	
HCT-116	> -4.00	
HCT-15	> -4.00	
HT29	> -4.00	
KM12	> -4.00	
KM20L2	> -4.00	
SW-620	> -4.00	
CNS Cancer		
SF-268	> -4.00	
SF-295	> -4.00	
SF-539	> -4.00	
SNB-19	> -4.00	
SNB-78	> -4.00	
U251	> -4.00	
XF 498	> -4.00	i
Melanoma		
LOX IMVI	> -4.00	
MALME-3M	> -4.00	!
M14	> -4.00	1
M19-MEL	> -4.00	
SK-MEL-2	> -4.00	i
SK-MEL-5	> -4.00	
UACC-257	> -4.00	Į.
UACC-62	> -4.00	
Ovarian Cancer		
IGROVI	> -4.00	
OVCAR-3	> -4.00	
OVCAR-4	> -4.00	
OVCAR-5	> -4.00	
OVCAR-8	> -4.00	}
Renal Cancer		
786-0	> -4.00	'
A498	> -4.00	
ACHN	> -4.00	
CAKI-I	> -4.00	
RXF 393	> -4.00	
RXF-631	> -4.00	
SN12C	> -4.00	
UO-31	> -4.00	
MG_MID	-4.00	
Delta	0.06	,
Range	0.06	
·/m·2c	0.00	
	+3	+2 +1 0 -1 -2 -3
	1 73	·~ T1 U -1 -4 •3

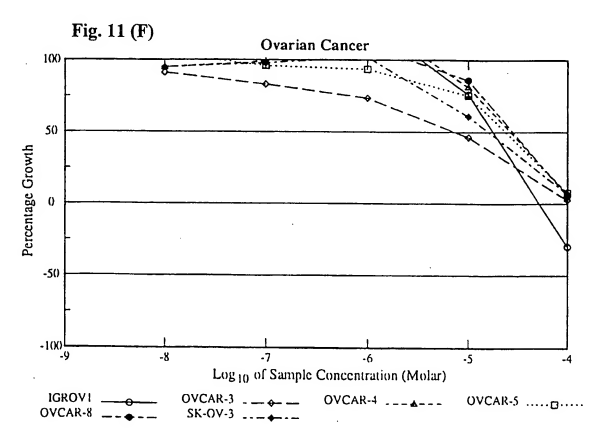


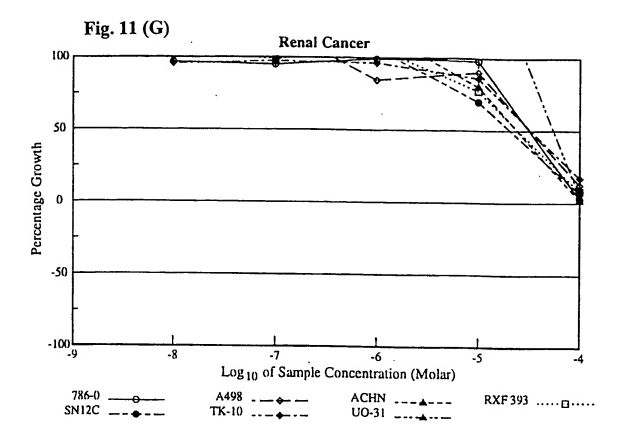


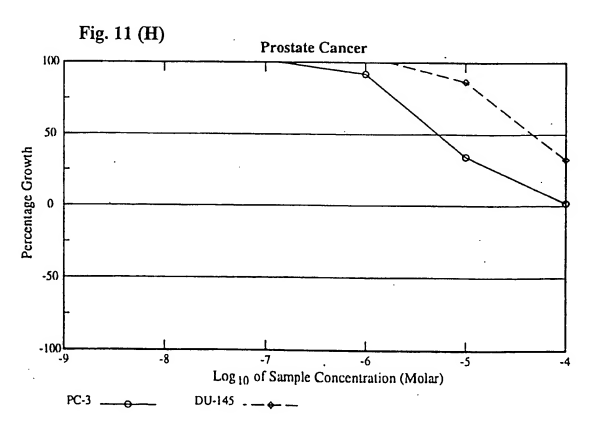


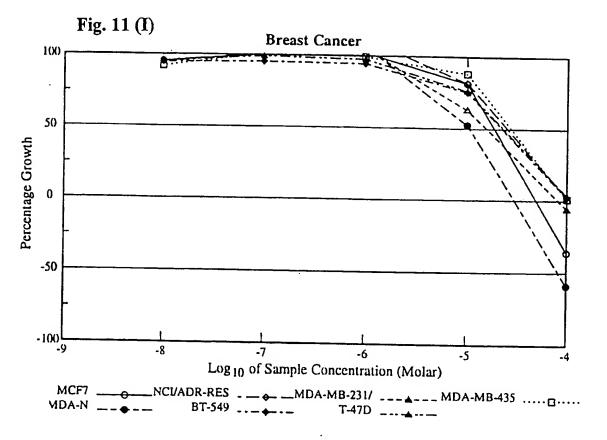






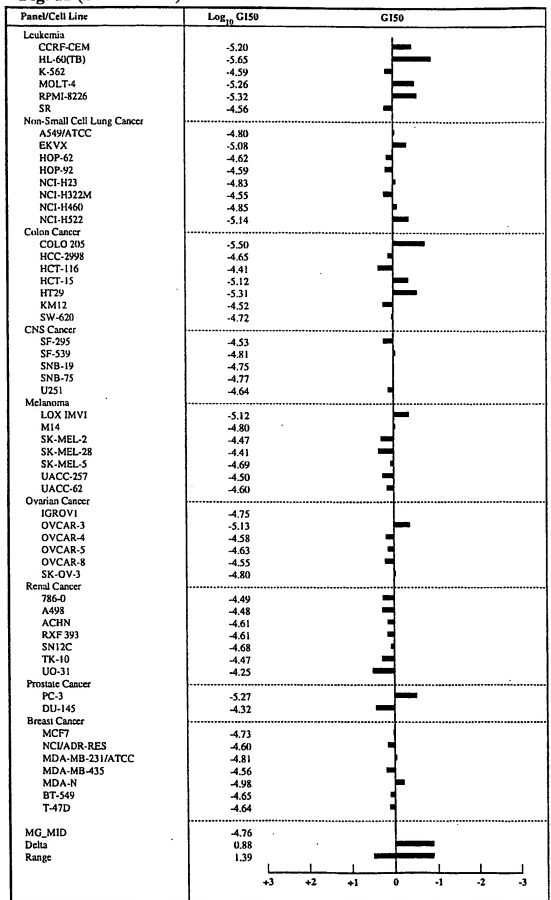






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Fig. 12 (Sheet 1 of 3)



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Fig. 12 (Sheet 2 of 3)

Fig. 12 (Sheet 2 01 3)	70/7/	
Panel/Cell Line	Log ₁₀ TGI	TGI
Leukemia		
CCRF-CEM	> -4.00	1
HL-60(TB)	> -4.00	1
K-562	> -4.00	
MOLT-4	> -4.00	
RPMI-8226	> -4.00	
SR	> -4.00	1
Non-Small Cell Lung Cancer		***************************************
A549/ATCC	> -4.00	1
EKVX	> -4.00	*
HOP-62	> -4.00	L_0.50
HOP-92	> -4.00	1
NCI-H23	> -4.00	1
NCI-H322M NCI-H460	> -4.00	<u> </u>
NCI-H400 NCI-H522	-4.30 4.30	
Colon Cancer	-4.38	
COLO 205	> -4.00	
HCC-2998	> -4.00 -4.34	_
HCT-116	-4.34 > -4.00	_
HCT-115	> -4.00 > -4.00	j .
HT29	> -4.00	
KM12	-4.15	l l
SW-620	> -4.00	[
CNS Cancer	- 1,04	
SF-295	> -4.00	
. SF-539	-4.41	
SNB-19	> -4.00	. •
SNB-75	> -4.00	•
U251	> -4.00	ą į
Melanoma	***************************************	
LOX IMVI	> -4.00	4
M14	-4.40	
SK-MEL-2	> -4.00	•
SK-MEL-28	> -4.00	4
SK-MEL-5	> -4.00	•
UACC-257	> -4.00	•
UACC-62	> -4.00	•
Ovarian Cancer		***************************************
IGROVI OVCAR-3	-4.28	
OVCAR-3	> -4.00	3
OVCAR-5	> -4.00 > -4.00	3
OVCAR-8	> -4.00]
SK-OV-3	> -4.00	3
Renal Cancer	~ ~1.U()	
786-0	> -4.00	***************************************
A498	> -4.00	1
ACHN	> -4.00	1
RXF 393	> -4.00	1
SN12C	> -4.00	
TK-10	> -4.00	1
UO-31	> -4.00	-
Prostate Cancer		
PC-3	> -4.00	4
DU-145	> -4.00	•
Breast Cancer		
MCF7	-4,31	–
NCI/ADR-RES	> -4.00	4
MDA-MB-231/ATCC	-4.09	1
MDA-MB-435	> -4.00	1
MDA-N	-4.53	· ·
BT-549	> -4.00	1
T-47D	> -4.00	1
NO ME	4	
MG_MID	-4.06	<u></u>
Delta	0.47	<u>-</u>
Range	0.53	, -
	.1	
	+3	+2 +1 0 -1 -2 -3
<u> </u>		

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Fig. 12 (Sheet 3 of 3)

Lcukemia CCRF-CEM HL-60(TB) K-562	> -4.00			2			
HL-60(TB)	> -4.00						
K-302	> -4.00			1			
MOLT-4	> -4.00						
RPMI-8226	> -4.00						
SR	> -4.00			1			
Non-Small Cell Lung Cancer						•••••	
A549/ATCC	> -4.00			4.0			
EKVX	> -4.00						
HOP-62	> -4.00						
HOP-92	> -4.00						
NCI-H23	> -4.00						
NCI-H322M	> -4.00						
NCI-H460	> -4.00						
NCI-H522	> -4.00						
Colon Cancer	> -4.00	***************************************					
COLO 205 HCC-2998	-4.02			1			
HCT-116	> -4.00			- [
HCT-15	> -4.00						
HT29	> -4.00			1			
KM12	> -4.00			1			
SW-620	> -4.00						
CNS Cancer				••••			
SF-295	> -4.00						
SF-539	-4.01						
SNB-19	> -4.00						
SNB-75	> -4.00			1			
U251	> -4.00						
Melanoma	> -4.00	,					
LOX IMVI	> -4.00			1			
M14 SK-MEL-2	> -4.00						
SK-MEL-28	> -4.00			İ			
SK-MEL-5	> -4.00						
UACC-257	> -4.00			ı			
UACC-62	> -4.00			1			
Ovarian Cancer					•••••	•••••	
IGROVI	> -4.00			1			
OVCAR-3	> -4.00		•				
OVCAR-4	> -4.00						
OVCAR-5	> -4.00			i			
OVCAR-8	> -4.00						•
SK-OV-3	> -4.00						
Renal Cancer	> -4.00						
786-0 A498	> -4.00						
ACHN	> -4.00						
RXF 393	> -4.00			-			
SN12C	> -4.00			1			
TK-10	> -4.00						
UO-31	> -4.00			1			
Prostate Cancer		,	•••••	••••			
PC-3	> -4.00						
DU-145	> -4.00						•
Breast Cancer			**********		•		
MCF7	> -4.00						1
NCI/ADR-RES	> -4.00						
MDA-MB-231/ATCC	> -4.00			l			
MDA-MB-435	> -4.00			L			
MDA-N	-4.08 > -4.00			Γ			
BT-549	> -4.00			1			
T-47D	> -4.00		*************			*******	
MG_MID	-4,00						
MG_MID Delta	0.08			þ			
Range	0.08			þ			
					1		لــــا
	i i	+3 +2	+1	0	-1	-2	-3

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- (57) Abstract

Certain derivatives of acridones and benzothiadiazines have been found to have anti-cancer properties by virtue of their specific inhibition of the cyclin D dependant kinase CDK4. These molecules inhibit CDK4 activity more than they inhibit the activity of other such kinases (e.g. CDC2 and CDK2). This specificity results in an improved therapeutic index when used as drugs to treat susceptible cancers.

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EE	Estonia	LR	Liberia	SG	Singapore		

Interna and Application No PCT/US 98/08602

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C07D219/08 A61K A61K31/435 A61K31/54 CO7D219/06 C07D219/04 C07D285/24 C07D285/16 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 CO7D A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ' Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Α BALL K L ET AL: "CELL-CYCLE ARREST AND 1-57 INHIBITION OF CDK4 ACTIVITY BY SMALL PEPTIDES BASED ON THE CARBOXY-TERMINAL DOMAIN OF P21WAF1" CURRENT BIOLOGY, vol. 7, no. 1, $\hat{1}$ January 1997, pages 71-80, XP002039815 X FR 2 676 737 A (GROUPE ENSEIGNEMENT RECH 1-21CHIM) 27 November 1992 40-43 see the whole document -/--Further documents are listed in the continuation of box C. X Patent family members are listed in annex. Special categories of cited documents : To later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. other means *P* document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 2 8. 05, 99 26 February 1999 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016 Steendijk, M

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	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
(MANNANI R ET AL: "Preparation of 3,3'-linked bis(acridinones), 9,9'-linked bis(thioacridines) and 3-3', 9-9' bi-linked bis(thioacridinones)" EUR. J. MED. CHEM. (EJMCA5,02235234);91; VOL.26 (1); PP.117-19, XP002075295 UFR PHARM.;GROUPE ENSEIGN. RECH. CHIM. THER. ORG. PHYS.; MARSEILLE; 13385; FR. (FR) see the whole document	
	MANNANI R ET AL: "Synthesis of two novel thioacridine derivatives and comparison of their in vitro biological activities" CHEM.—BIOL. INTERACT. (CBINA8,00092797);90; VOL.74 (3); PP.291—303, XP002075296 FAC. PHARM.;GROUPE ETUD. RECH. CHIM. THER. ORG. PHYS.; MARSEILLE; FR. (FR) see the whole document	1
X	CHEMICAL ABSTRACTS, vol. 112, no. 19, 7 May 1990 Columbus, Ohio, US; abstract no. 178637g, MARTYNOVSKII ET AL.: "Synthesis, physicochemical properties and pharmacological activity of carbonyl-substituted 2-methoxy-9-thioacridines" XP002091482 & Izv. Timiryazevsk. S-kh. Akad., 1989, (5), 174-181 see abstract	1-20,25
A	CHEMICAL ABSTRACTS, vol. 070, no. 11, 17 March 1969 Columbus, Ohio, US; abstract no. 047270, WYSOCKA-SKRZELA B ET AL: "Tumor inhibiting compounds. XLIV. Syntheses of N-substituted 1-, 2-, 3-, and 4-methoxyacridones and thioacridones. 2" XP002075299 see abstract & ROCZ. CHEM. (ROCHAC);68; VOL.42 (10); PP.1755-61, POLSKA AKAD. NAUK;GDANSK; POLAND/	1-57
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ation) DOCUMENTS CONSIDERED TO BE RELEVANT	<u> </u>
Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
TAIT A ET AL: "Antitumor activity of methyl-4H-1,2,4-benzothiadiazin-3-yl-carba mod ithioate-S,S-dioxide" BOLL. CHIM. FARM. (BCFAAI,00066648);90; VOL.129 (9); PP.273-5, XP002075297 UNIV. MODENA:DIP. SCI. FARM. ITALY (IT)	2-14,39, 40
see the whole document	1,15-38, 41-57
CHEMICAL ABSTRACTS, vol. 124, no. 17, 22 April 1996 Columbus, Ohio, US; abstract no. 232400u, PARK ET AL.: "Synthesis and cytotoxic properties of 3,4-dihydro-3-oxo-2H-1,2,4-benzothiadiazin e 1,1-dioxides"	2-14,39, 40
& Yakhak Hoechi, 1995, 39(6), 631-635 see abstract	1,15-38, 41-57
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CHEMICAL ABSTRACTS, vol. 120, no. 25, 20 June 1994 Columbus, Ohio, US; abstract no. 315309, LEE ET AL.: "Inhibition of cell growth by K+ channel modulators is due to interference with agonist-induced Ca2+release" XP002091483 & Cell. Signalling, 1993, 5(6), 803-809 see abstract	2-14,39, 40
CHEMICAL ABSTRACTS, vol. 73, no. 15, 12 October 1970 Columbus, Ohio, US; abstract no. 75517, NAKAHARA ET AL.: "Oncostatic activities of some fluoro compounds against Ehrlich carcinoma in mice" XP002091484 & Wakayama Daigaku Gakugeigakubu Kiyo, Shizen kagaku 1968, no. 18, 15-17 see abstract	2-14,39, 40
	TAIT A ET AL: "Antitumor activity of methyl-4H-1,2,4-benzothiadiazin-3-yl-carba mod ithioate-S,S-dioxide" BOLL. CHIM. FARM. (BCFAAI,00066648);90; VOL.129 (9); PP.273-5, XP002075297 UNIV. MODENA;DIP. SCI. FARM.; ITALY (IT) see the whole document

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	Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.				
X	REDDY ET AL.: "Synthesis & biological activity of 3-pyrazolyl-4-substituted-2H-1,2,4-benzoth iadiazine 1,1-dioxides" INDIAN JOURNAL OF CHEMISTRY, vol. 24B, 1985, pages 1295-1297, XP002091480 see the whole document	2-14,39, 40				
X	JIANG ET AL.: "Synthesis and biological evaluation of 2-styrylquinazolin-4(3H)-ones, a new class of antimitotic anticancer agents which inhibit tubulin polymerization" J.MED.CHEM., vol. 33, 1990, pages 1721-1728, XP002091481 see page 1723; table IV	2-14,39, 40				
X	CHEMICAL ABSTRACTS, vol. 113, no. 13, 24 September 1990 Columbus, Ohio, US; abstract no. 115267, SADANA ETAL.: "Antibacterial activity and synthesis of 4-(substituted benzylamino)-1,2,3-benzothiadiazine 1,1-dioxides" XP002091485 & Indian J. Chem., Sect. B, 1990, 29b(6), 598-599. RN: 129226-06-8 see abstract	2-20, 36-38				
X	US 3 090 783 A (YALE) 21 May 1963 see the whole document	2-20,29				

International application No. PCT/US 98/08602

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest X The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1,21,28,43,50 and (part) 2-20, 22-27,29-42,44-49, 51-57

Compounds, antineoplastic compositions and methods involving at least one thioacridone, possibly in combination with a benzothiadiazine.

2. Claims: 2-20, 22-27,29-42,44-49,51-57 (all part)

Antineoplastic compositions and methods involving a benzothiadiazine, not in combination with thioacridinones

Information on patent family members

PCT/US 98/08602

	tent document in search repor	t	Publication date	Patent family member(s)	Publication date
FR	2676737	Α	27-11-1992	NONE	_
DE	3331459	Α	01-03-1984	NONE	***************************************
WO	9749692	Α	31-12-1997	AU 3166497 A	14-01-1998
US	3090783	Α	21-05-1963	NONE	